# MANGANESE SULPHUR INTERACTIONS IN IRON CARBON ALLOYS

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In Partial Fulfilment of the Requirements
for the Degree of

DOCTOR OF PHILOSOPHY

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to the

DEPARTMENT OF METALLURGICAL ENGINEERING
INDIAN INSTITUTE OF TECHNOLOGY, KANPUR
SEPTEMBER, 1984



#### CERTIFICATE

Certified that the work reported in this thesis entitled 'Manganese Sulphur Interactions in Iron-Carbon Alloys' by A.K. Pattjoshi has been carried out under my supervision and has not been submitted elsewhere for the award of degree.

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A. K. Palljeshi

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#### SYNOPSIS

MANGANESE-SULPHUR INTERACTIONS IN IRON CARBON ALLOYS

A Thesis Submitted in Partial Fulfilment of the Requirements for the Degree of

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by

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Department of Metallurgical Engineering Indian Institute of Technology, Kanpur September, 1984

Studies on sulphur control in iron making and steel making processes or by means of external treatment of metal in the ladle have gained importance in recent years to meet challanges for production of low sulphur steel at lower cost for newer and newer applications in space, ocean, defence Basic studies are important in understanding the mechanism of sulphur removal from the metal to slag. Thermodynamics of sulphur reactions and desulphurising powers of various blast furnace and steel making slags have been investigated by many workers by equilibriating the melt with known H2/H2S ratio or with a known quantity and composition of slag at a fixed temperature. present work attempts are made to study the effect of Mn on the desulphurisation of iron-carbon melts due to formation of MnS with and without addition of some oxide slags of CaO-SiO<sub>2</sub>-Al<sub>2</sub>O<sub>3</sub> system in the temperature range of 1200°-1400°C. Such temperatures usually occur in the transit of

the hot metal from the blast furnace to the steel making unit and in presence of Mn, sulphur drops are expected to occur.

Laboratory experiments included fabricating a silicon carbide furnace, making iron-carbon alloys, melting the alloy under argon atmosphere in a graphite crucible, drawing the samples carefully with a quartz tube and the analysis of sulphur and manganese in the laboratory.

The effect of Mn in metal on the desulphurising power of oxide slags was studied by making synthetic slags of CaO-SiO2-Al2O3 in the laboratory and equilibrating the melt at 1400°C with those slags for 4-6 hours of duration. Samples were either drawn from the melt or taken from the solidified mass in the crucible after the duration of experiment. Pure lime was also used as a desulphuriser in some experiments for comparison purposes. Reliability of data was checked by repeating a few experiment, by making more than one analysis of some samples and by calibrating the apparatus with a standard steel sample supplied by British chemical standards. Thermocouples were suitably calibrated against standard thermocouples and with melting point of pure copper. Raw materials were characterised by X-ray diffraction and by chemical analysis in the laboratory.

Experiments were done by stepwise addition of manganese to the melt of Fe-C-S at constant temperature or by cooling and heating of Fe-C-Mn-S melt. The solubility

of sulphur determined experimentally at lower temperatures of 1200°C matched well with the reported data of Morris\*, where as at 1400°C the sulphur levels in the melt were found to be less in the present investigation.

A regular solution model has been proposed for finding the activity coefficient of sulphur at temperatures lower than 1600°C as data existed in the literature only at 1600°C.

The calculated values of the sulphur content in carbon saturated metal at different temperatures and manganese contents of metal were compared with the experimental results. Good agreement was reached in most of the cases if the activities of MnS and FeS in slag were taken to be the same as their weight fractions and a discrepancy of -1.0 to -1.8 kcal/mole in the reported standard free energy of formation of MnS was allowed. These limits are within the accuracy of the reported data.

From the slag metal equilibrium studies at 1400°C, manganese up to 2 pct. was found to decrease the sulphur content of metal at all combinations of slag volume (10 to 20 pct.) and slag basicity (CaO/SiO<sub>2</sub> ratio of 1 to 2). A multicomponent multiphase mathematical model was developed to analyse the experimental findings. Sulphur partition ratios were calculated both from the model and the experimental data. Exact role of manganese in lowering the sulphur content of the iron-carbon alloy melt could not be identified. More

<sup>\*</sup> J.P. Morris, J. of Metals, (September 1952), p. 939-940.

scientific data: under the laboratory and plant condition are recommended to understand the mechanism of sulphur transfer from slag to metal in presence of manganese in the melt.

#### CHAPTER - I

#### INTRODUCTION

# 1.1 Effect of Sulphur on Mechanical Properties of Steel

Sulphur in steel is considered to be harmful, because the presence of sulphur lowers the mechanical properties of steel. Free cutting steel is the only grade of steel where the sulphur content is kept deliberately high, i.e. 0.22 weight pet or more. Iow sulphur in steel is essential for improved notch toughness, ductility, cold pressing properties and to avoid cracking failure as well as directional anisotropy in mechanical properties on hot working. For the production of quality steel and for super duty applications, the sulphur in steel should be generally less than 0.01 percent. It is generally specified that the final product i.e. steel should have 0.025-0.040 percent sulphur for general applications and the modern trend is towards lower and lower sulphur contents in steel.

Some of the detrimental effects of sulphur in steel are elaborated further below.

### 1.1.1 Ductility of steel

A major factor determining the yield in hot working operations is the ductility of steel. Some steels having lower ductility restrict their range of working temperature. Elements which form low melting point compounds or inclusions

deform differentially leading to cracking or large scale fracture. Sulphur in steel is known to form a low melting point compound FeS and contributes to red shortness. This red shortness is manifested as a marked lack of ductility in the temperature range 850-1050°C when sulphur content is greater than 0.005 percent. The addition of manganese above a critical value, improves the ductility of the steels which are susceptible to red shortness. The stoichometric ratio of 'Mn' to 'S' in MnS is 1.72 and at ratio lower than this, some 'S' should be present in inclusions as FeS or (Fe, Mn)S. The loss of ductility increases with increasing amount of inclusion material.

## 1.1.2 Anisotropy

The sulfides are very plastic at high temperatures and they get elongated in the direction of rolling. They thus contribute to the anisotropy of mechanical properties relative to the longitudinal and transverse directions. Sulfides lower the ductile fracture level of the steel longitudinally as well as transversely.

#### 1.1.3 <u>Impact</u>

In the case of nickel maraging steels, the influence of sulphur on impact properties is highly pronounced and maximum of 0.005 to 0.01 pct, of sulphur can be allowed in such steels. In high strength steels even lower sulphur contents have a marked influence on resistance to sudden crack propagation in the longitudinal as well as in the transverse direction.

#### 1.1.4 Fatigue

Depending upon the steel composition sulphur lowers its transverse fatigue resistance. An increase in sulphur content of steel has detrimental effect on transverse fatigue resistance.

## 1.1.5 Weld defects

Lamellar tearing is considered to be the most serious cracking problem during welding of structural and pressure vessel steels. Sulphur in steel gives rise to lamellar tearing in weld joints. Also sulphur imparts porosity in weld joints. The concentration, size and distribution of sulphide inclusions are controlling factor for the problems in weld joints.

#### 1.1.6 Conclusion

It is clear from the foregoing sections that presence of sulphur in steel is not at all desirable. The sulphur levels in structural steels should be less than 0.035 Pct and less than 0.02 Pct for quality steels.

To achieve low sulphur in steel, varieties of processes are in operation throughout the world. The origin and control of sulphur in steel are dealt in the following sections.

## 1.2 Origin of Sulphur in Steel

Bulk of the steel is commercially produced through the traditional route of blast furnace/ID or blast furnace/ openhearth, while a small fraction is also produced by other methods such as electric arc furnace melting of scrap, DR/EAF route etc.

The major source of sulphur input to the blast furnace is through coke, with a very small fraction entering with sinter or iron ore. Sulphur in the blast furnace is favourably distributed between slag and metal due to the reducing conditions inherent in blast furnace smelting of pig iron. A typical sulphur balance in a blast furnace is shown in Table 1.1.

It is apparent that a very small fraction of sulphur entering with the burden leaves with flue gases. As coke burns at the tuyeres, sulphur is picked up by H<sub>2</sub> and CO gases to form H<sub>2</sub>S and COS. As these gases come in contact with iron bearing and CaO bearing materials, sulphur is absorbed to form FeS and CaS and it descends to the bottom along with the rest of materials in the molten form.

## 1.3 Control of Sulphur in Steel

# 1.3.1 Sulphur control in blast furnace iron

Sulphur in the charge materials is absorbed by the metal and slag in the bosh or shaft and the final transfer of sulphur from metal to slag occurs in the hearth.

If 'a' the distribution coefficient of sulphur is defined as ratio of wt pct 'S', in slag to wt pct 'S' in metal i.e.

$$\alpha = \frac{(\text{Pct.S})_{\text{slag}}}{[\text{Pct.S}]_{\text{metal}}}$$
(1.1)

'S' level in hot metal can be written as follows:

$$[S] = \frac{100 \, S_{\text{input}}}{1000 + \alpha \, W_{\text{elec}}} \tag{1.2}$$

where Sinput and Wslag are the weights of sulphur entering with burden, and slag respectively in kg/THM. It is assumed

Table 1.1 (Ref.1)

A Typical Blast Furnace Sulphur Balance

Material	Sulphur input			Sulphur output	
	Kg/IHM	Pct. of fotal	Kg/THM	Pet. of Total	
Coke	6.0	80	ann and a 1871 a		
Charge Materials	1.5	20			
Iron	-	: :/:-	0.30	4.0	
Flue dust and gases	-	<del>-</del>	0.37	5.0	
Slag	<del>.</del>		6.83	91.0	
Total	7.5	ood), deestis in viintallik, elklustuurse myök, mel	7.5-	ang unger and under various strands und Ethanh a	

that no sulphur leaves with flue gases. Equation (1.2) is plotted in Figure 1.1. Iow sulphur in hot metal can be achieved only at higher 'a' values unless slag volumes are excessively high. High slag rates can lead to high coke rates which in turn will increase Sinput values and hence [S]. Hence the only effective method of producing low[S] iron is to adjust slag composition and temperature in such a manner as to get high sulphur distribution ratios in the blast furnace.

High sulphur distribution ratios are obtained in a blast furnace by maintaining high slag basicity i.e. (CaO+MgO)/SiO<sub>2</sub> ratio and high metal temperature and their effects will be discussed later. These factors lead to an increase in carbon rate of the furnace as shown by Flint<sup>2</sup> by the regression analysis of large plant data. He reported that carbon rate increases in a curvilinear fashion as shown in Figure 1.2 and this data is still used by many operators in making the techno economic studies to control sulphur in the blast furnaces.

## 1.3.2 Sulphur control in steel making

Removal of sulphur from the hot metal during refining by the oxidation process is very limited due to high FeO content of the slag. It is not possible to have sulphur distribution coefficient values greater than 5 or 10 and it puts the limit to which sulphur can be removed in a steel making process. In most of cases only fifty percent of sulphur can be removed from the melt during refining. Favourable conditions for sulphur removal include high basicity i.e. (CaO+MgO/SiO2 ratio,

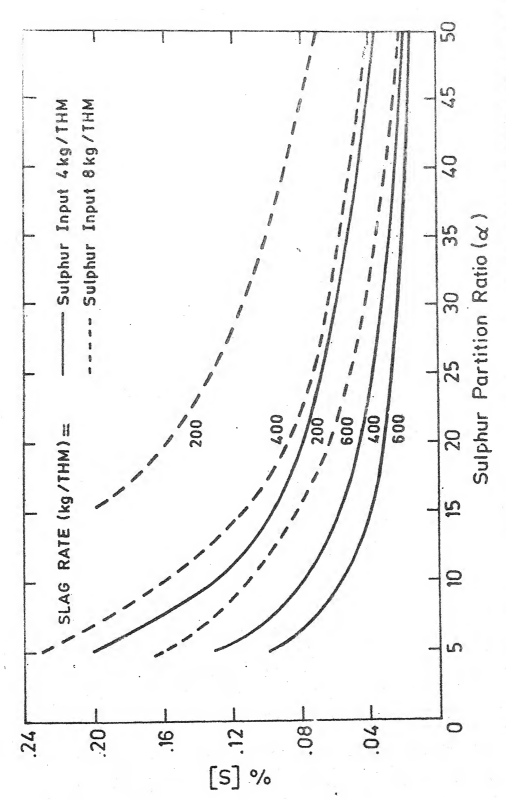


FIG. 1.1 SULPHUR LEVELS IN HOT METAL VERSUS SULPHUR PARTITION RATIO IN BLAST FURNACE.

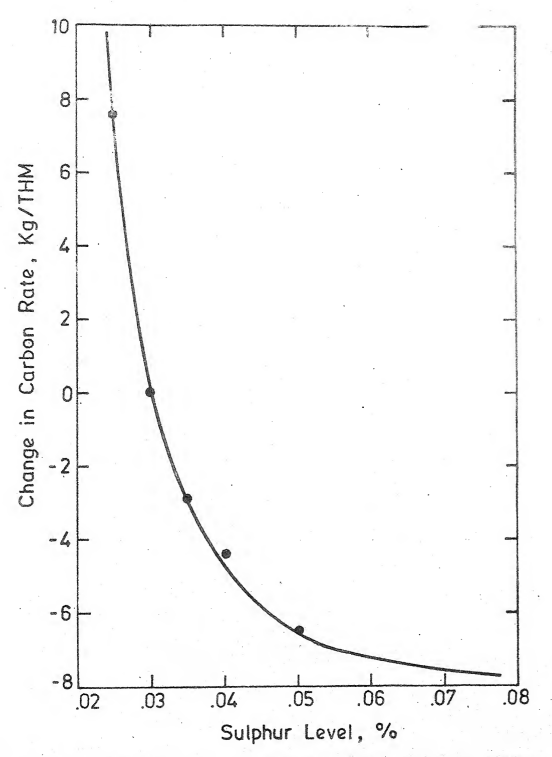


FIG. 1.2 EFFECT OF SULPHUR LEVEL IN HOT METAL ON CARBON RATE IN BLAST FURNACE. (Ref. 2)

good fluidity of slag and high temperature. Sulphur input via hot metal, ores, fluxes and fuels must be minimised to produce low sulphur steel for achieving better mechanical properties. Typical sulphur balances in open hearth and L.D. processes of steel making are given in Table 1.2.

Electric furnaces offer ideal conditions for producing very low sulphur levels because high sulphur distribution ratios are possible in the reducing period, where FeO in the slag may not exceed 0.1 percent. Full discussions on the thermodynamics and kinetics aspects of sulphur reactions during steel making lie outside the scope of present investigation and are not reported in this work.

## 1.3.3 External desulphurisation of iron

External treatment of hot metal to get rid of sulphur is being widely practised all over the world. The technoceconomic considerations do not normally justify the sulphur removal to low levels in either blast furnace iron making or in LD steel making. Desulphurising agents such as soda ash, lime, calcium carbide or magnesium are either added or injected to the metal in a suitable manner to enhance the rate of sulphur removal from the metal to the slag. The slag formed must be removed from the ladle more effectively otherwise sulphur reversion may occur during steel making. The following factors will govern the choice of the process technology and the dasulphurising agents in any steel plant.

- (1) Hot metal composition and its temperature.
- (2) Initial and the final sulphur levels in hot metal.

Table 1.2

A Typical 'S' Balance in LD/BOH Steel Making

Process	Input,	Kg/THM	Output,	Kg/THM
ogi, in 7-86° (48° -7-monostourily 40°7-64° (international money transposition international continues court	Sin	Sin		out
	Hot metal	0.48	Steel	0.35
LD	Scrap	0.07	Slag	0.20
	Total	0.55	Total	0.55
	Hot metal	0.13	Steel	0.250
	Scrap	0.15	Slag	0.135
ВОН	Time stone and dolomite	0.06		
	Fluorspar	0.020		
	Flame gasses	0.025		
	Total	0.385	Total	0.385

- (3) Space limitations in the plant.
- (4) Cost of installation of desulphurisation unit and eperation costs.
- (5) Effectiveness of slag removal after desulphurisation.
- (6) Availability and cost of desulphurising agents.

The following difficulties may be encountered during any external desulphurisation:

- freezing of the top layer of metal in the laile may occur to give high skull losses. This also makes addition of desulphurising agents to the bath very difficult.
- (2) Sulphur rich blast furnace slags may accompany the metal and create difficult conditions for 'S' removal.
- (3) Ladle lining may get damaged due to addition of some desulphurising agents.
- (4) Environmental problems.
- (5) Extra handling and delay in the transport of the iron ladles.
- (6) Inefficient removal of slag from the ladles after the external desulphurisation of hot metal.

Various methods and desulphurising agents have been used in the last two decades and those are summarized in Table 1.3.

The station for ladle treatment of metal can be located either near the blast furnace or near the mixer in the steel making shop depending upon the existing conditions in a

Table 1.3
Methods of Desulphurisation

Desulphu- rising agents	Mode of addition	Reagent Quantity Kg/THM	'S'-Range Pct.	Tempera- ture drop °C	Remarks
Soda ash	Bulk addition	10	.10035	60–70	Attacks ladle lining,forms corrosive liquid slag
Inme	Injection	10-20	.0502	70–100	Large slag volume
Time - Soda ash	Injection	8–10	.0502	50 <b>–</b> 80	
Calcium Carbide	Injection	4	,0502	15-20	Expensive, good for low sulphur
Mg-Coke (45 percent Mg)	Plunging	0.7	.048022	5–15	
Mg-granu- les	Plunging	.7-1.2	.05005	5–10	Good for very low sulphur

particular plant. It may also be possible to desulphurise the metal externally in the blast furnace cast house itself where slag can be removed efficiently in the runner but initial sulphur will not be known and the process may create practical troubles for the tapping crew. In most cases a separate external desulphurisation station is installed between the blast furnace and mixer for taking all necessary steps and precautions for the process optimisation.

#### 1.4 Scope of Present Investigation

The present investigation was undertaken primarily to study the manganese/sulphur interactions in iron carbon alloys in the temperature range of  $1200^{\circ}$  -  $1400^{\circ}$ C. The role of manganese on the desulphurising power of oxide slags at  $1400^{\circ}$ C was also studied as this information was found to be lacking in the literature. Suitable arrangements were made to design and conduct a few experiments under controlled conditions. The experimental results are supplemented by theoretical calculations based on the available thermodynamic data relevant to the systems of interest in the present study.

The comprehensive literature review is given in the next chapter which is followed by full details on the plans of the present work. Results of the theoretical calculations are given in Chapter 4 while Chapters 5 and 6 give experimental details and results respectively. Discussion on the findings of the present work and comparison with the results of other investigators are given in Chapter 7. Chapter 8 gives the summary

and conclusions of this work while recommendation for further work are given in Chapter 9. A complete biblography is given at the end of this thesis.

#### CHAPTER - II

#### LITERATURE REVIEW

#### 2.1 Introduction

Scientific investigations on desulphurisation of pig iron started as early as the 1930's and since then much work has been done to study the slag-metal and gas-metal equilibria. Some fundamental work pertinent to desulphurisation of iron has also been carried out. A comprehensive literature survey is presented in this chapter.

#### 2.2 Preliminary Studies

First studies on the desulphurisation of pig iron were done by Wentrup<sup>3</sup>, Holbrook and Joseph<sup>4,5</sup> and Oelsen<sup>6</sup>.

Wentrup found that desulphurisation was improved by lowering the temperature in presence of manganese in pig iron. For the Fe-S-Ca-O-Mn-Si system, Wentrup defined the desulphurisation ratio i.e. the distribution of sulphur between metal and slag by the ratio of Total(S) in slag to[S] in metal. The values of Slmetal was experimentally found to be 876.5 for slag containing 0.1 pct FeO, 30 pct CaO and zero pct SiO<sub>2</sub> and 145.2 for slag containing 2.4 wt pct FeO.

Wentrup concluded that if FeO and SiO<sub>2</sub> content of slag were kept low, then lime in slag could be an effective desulphuriser.

Holbrook and Joseph<sup>4</sup> studied the desulphurising power of blast furnace slags, and defined it as the ratio

Their experiments were carried out under standardized laboratory conditions and revealed that, in magnesia free slags the desulphurising power increased rapidly as lime replaced alumina and slowly as alumina replaced silica. The desulphurising power decreased slightly with the addition of magnesia to the acidic slags and increased markedly when magnesia was added to very basic slags.

Systematic work on the distribution of sulphur between slag and metal by first finding the activity of sulphur in the iron base alloys was carried out by Chipman and co-workers7. Their findings are described in the following section.

#### 2.3 Activity of Sulphur in Tiquid Iron Alloys

#### Experimental approach to the determination of 2.3.1 thermodynamic data

The experimental approach to determine the activity of sulphur in the melt required the attainment of equilibrium between the sulphur in the melt and gaseous atmospheres containing hydrogen and hydrogen sulphide. The reaction and its equilibrium constant are written as follows:

$$\underline{S} + H_2(gas) = H_2S (gas) \tag{2.1}$$

The standard free energy change of reaction (2.1) can be written as

$$\Delta_{2.1}^{G_{2.1}} = - \underset{p_{H_2}}{\text{RT ln } K_{2.1}} \tag{2.2}$$

$$K_{2.1} = p_{H_2} a_s \tag{2.3}$$
or  $K_{2.1} = \frac{p_{H_2}}{p_{H_2}} \times \frac{1}{f_s [\text{wt% s}]}$ 

or 
$$K_{2.1} = \frac{p_{H2S}}{p_{H2}} \times \frac{1}{f_s[wt\% s]}$$
 (2.4)

where,  $p_{H_2S}$  and  $p_{H_2}$  are the partial pressures of  $H_2S$  and  $H_2$  respectively in the gas mixture,  $f_s$  is the activity coefficient of sulphur in the liquid metal and [wt % S] is the percentage of sulphur in the melt.

The activity of sulphur, as, in the melt can be described by the following equation;

$$a_{s} = f_{s} [wt \% S]$$
 (2.5)

In a multi component system containing solutes 1,2,3,4... log  $f_s$  may be represented by the following equation;

$$\log f_s = \log f_{s_1} + \log f_{s_2} + \dots$$
 (2.6)

where, 
$$\log f_{si} = e_s^i [wt \% i]$$
 (2.7)

 $e_s^i$  is the interaction parameter (Henrian) i.e. the interaction of sulphur with solute 'i' and is defined as  $e_s^i = \frac{\partial \log f_{si}}{\partial \left[wt/s \ i\right]}$  where,  $f_s \to 1$  as all the wt of solutes 1,2,3 — +0. Hence at zero concentration of sulphur, the equation (2.4) reduces to the following;

$$K_{2.1} = \frac{p_{H_2S}}{p_{H_2}} \times \frac{1}{[wt \% S]}$$
 (2.8)

The R.H.S. term of equation (2.8) may be described as  $K_{2.1}$  at any other concentration of sulphur. By fixing  $p_{H2S}/p_{H2}$  ratios and finding the corresponding equilibrium [ wt g S] in the melt,  $K_{2.1}$  values can be computed. A typical plot of  $\log K_{2.1}$  versus sulphur percent is shown in Fig. 2.1. By extrapolating  $K_{2.1}$  to zero pct sulphur, value of  $K_{2.1}$  can be determined. Then for any finite concentration, activity coefficient of sulphur can be determined as follows;

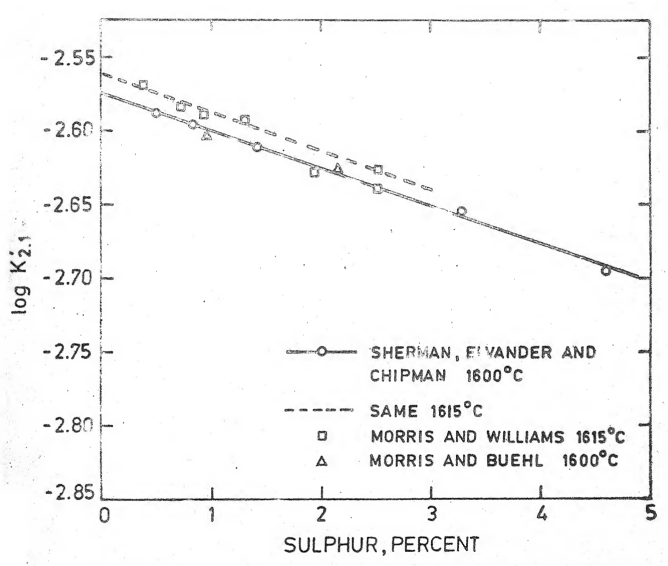


FIG. 2.1 EFFECT OF SULPHUR CONCENTRATION ON K'2.1. (Ref. 8,9,1

$$f_s = \frac{K_{2.1}^{\prime}}{K_{2.1}} \tag{2.9}$$

Similarly activity co-efficient f<sub>si</sub> in presence of solute 'i' in the melt can be determined and from equation (2.7) corresponding interaction parameter can be calculated.

### 2.3.2 Fe-S system

The standard free energy of formation of  $H_2S$  from gases such as  $H_2$  and  $S_2$  was reported as follows  $^{11}$ ;

$$H_2 + \frac{1}{2} S_2 = H_2 S_1 \Delta G_{2.10}^0 = -21820 + 11.92 \text{ T Cal.}$$
 (2.10)

Ban-ya and Chipman<sup>12</sup> calculated the standard free energy of formation of reaction (2.10) by finding the heats and entropies of formation and reported the following;

$$\Delta G_{2.10}^{0} = -21,530 + 11.73 \text{ T Cal.}$$
 (2.11)

They also determined the equilibrium constant experimentally as function of temperature in the range of 1500-1600°C and reported the results as follows;

$$H_2 + \underline{S} = H_2 S \tag{2.12}$$

$$\log K_{2.12} = -\frac{2350}{T} - 1.34 \tag{2.13}$$

or 
$$\Delta G_{2.12}^{0} = 10750 + 6.13 \text{ T Cal.}$$
 (2.14)

Combining equations (2.11) and (2.14);

$$\frac{1}{2}S_2 = \frac{S}{2}, \Delta G_{2.15}^0 = -32,280 + 5.60 \text{ T Cal.}$$
 (2.15)

Heat of solution of 'S' in iron was estimated as -32.28 KCal/mole with uncertainty of ± 2.5 Kcal/mole at 1600°C, but corresponding

uncertainty in AGO did not exceed 0.4 kcal.

Activity coefficient 'fs' varied with 'S' content of iron at 1600°C up to 3 pct 'S' as follows 12;

$$log f_s = -0.0282[\% S]$$
 (2.16)

From this  $e_s^s$  is calculated, -0.0282 which agrees with the value used by investigators earlier<sup>7</sup>.

## 2.3.3 Fe-C-S system

The solubility of sulphur in iron-carbon alloys was investigated by Morris and Buehl $^{10}$  by equilibrating the melt with  $\rm H_2$  -  $\rm H_2S$  gas mixtures as explained earlier. The results of Morris and Buehl are given in Fig. 2.2.

Increase in the activity coefficient of sulphur due to presence of carbon in the carbon saturated melts was found to be much greater than the earlier studies reported by Kitchner et al. 13, whose dafa could not be relied upon due to their wide scatter.

From Fig. 2.2,  $e_s^c$  value could be 0.12 upto carbon concentrations of 1 or 2 wt pct, but for carbon concentration beyond 2 pct,  $\log f_s$  versus [wt % C] plot is not linear and equation 2.7 can not be applied directly to find  $e_s^c$  value. The slope of  $\log f_s$  versus [% C] plot at 4-5 pct carbon level is around 0.24 which has been reported as the interaction parameter value by some investigators later  $^{31}$ .

Some investigators have reported an average value of  $e_s^c$  as 0.15 for thermodynamic calculation purposes to give the same value of  $f_s$  as determined experimentally by Morris and Buehl 10

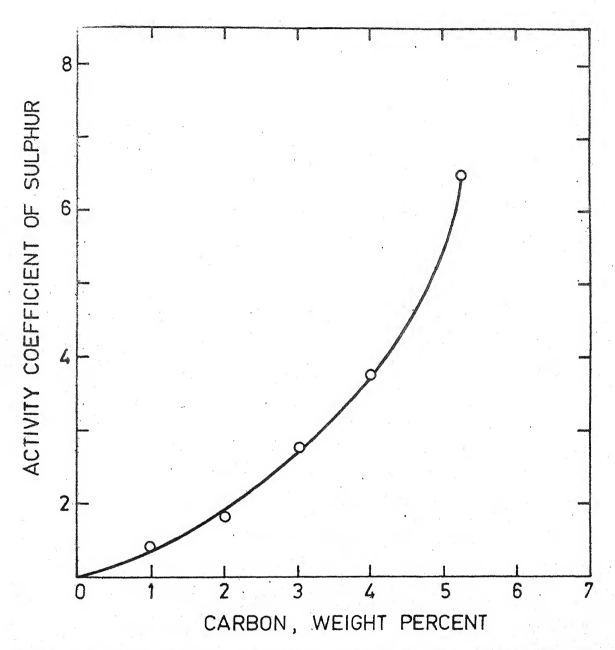


FIG. 2.2 EFFECT OF CARBON CONTENT ON ACTIVITY CO-EFFICIENT OF SULPHUR IN IRON AT 1600°C. (Ref. 10)

in the carbon level of 3-5 pct. It might be desirable to use 4.35 pct. Carbon in iron as its standard state as mentioned by Morris and Buehl, who reported the following;

$$\frac{1}{2} S_2(g) = \underline{S} (4.35\% C) \qquad (2.17)$$

$$\Delta G_{2.17}^{0} = -23,880 + 3.90 \text{ T Cal.}$$
 (2.18)

Equation (2.17) is related to the equation (2.15) by the following;

$$\Delta G^{0}_{(4.35\% C)} = \Delta G^{0} + RT \ln f_{s}^{C}$$
 (2.19)

 $f_s^c$  is found to be 4.2 at  $1600^\circ$ C at 4.35% C. Since the slope of  $log f_s$  versus [%C] plot is 0.24 at this carbon level,  $f_s^c$  at any other level of carbon can be determined as follows;

$$\log f_s^c = \log 4.2 + 0.24 (x - 4.35)$$
 (2.20)

where, x is the Wt. Pct Carbon in the melt.

# 2.3.4 Effect of alloying elements

Sherman and Chipman investigated activity of S in melt of iron containing various alloys such as Mn, P, Al, Cu, Si etc. at  $1600^{\circ}$ C. Their results are summarized in Figure 2.3. From  $\log f_{\rm S}^{\rm i}$  versus Wt% i plots in each case,  $e_{\rm S}^{\rm i}$  can be calculated applying equation (2.7). The interaction parameters value are summarized in Table 2.1.

Their results of Fe-S-Si system agreed with those obtained by Morris and Williams 9.

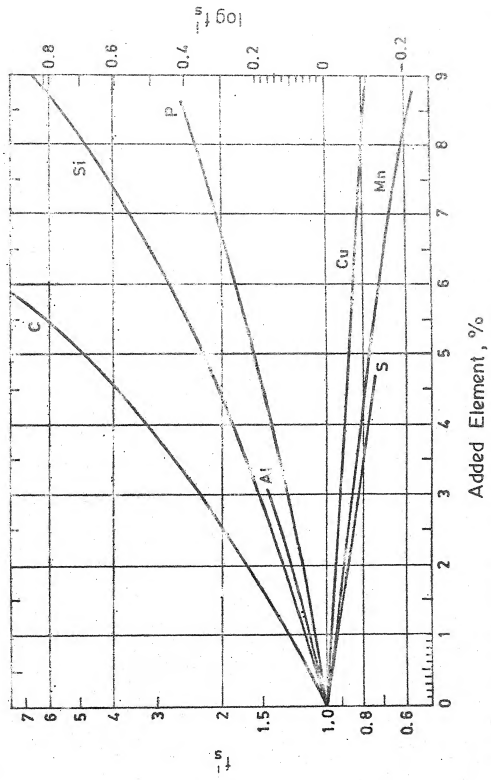


FIG. 2.3 EFFECT OF ALLOYING ELEMENTS ON ACTIVITY COEFFICIENT OF SULPHUR IN IRON AT 1600°C. (Ref. 7)

Table 2.1 (Ref.7)

Effects of Small Concentrations of Added Substances on Activity Coefficient of Sulphur

i = third component	$e_{S}^{i} = \partial \log f_{S}/\partial (\% i)$
C	+ 0.113
Şî	+ 0,065
P	+ 0.043
Al.	+ 0.054
Cu	- 0.013
${ t M}{ t n}$	- 0.025

Morris  $^{14}$  investigated the activity of sulphur in the melt containing carbon and manganese by equilibrating with a known  $H_2S/H_2$  ratio at  $1600^{\circ}$ C. His results are given in Table 2.2. His experimental results match well with those of Sherman and Chipman as can be seen by the good agreement in  $f_s$  values in Table 2.2.

# 2.4 Slag Metal Equilibrium

Hatch and Chipman<sup>15</sup> made an extensive scientific study of the sulphur distribution equilibria between blast furnace slag and metal. They melted 200 g of metal and 400 g of slag in an induction furnace, in a graphite cruicble under controlled atmosphere of carbon monoxide. Constant stirring of the batch was provided using a graphite rod.

Equilibrium was reported to have been reached in 4 to 6 hours of time depending on the viscosity, temperature and composition of the slag. Attainment of silicon equilibrium was reported slower than that of sulphur. Their equilibrium data are summarised in Fig. 2.4.

Their results are often used to determine the equilibrium sulphur distribution ratio (% S) /[% S] at different composition of the slag for comparison with actual values obtained in the iron making and steel making conditions. Calculated sulphur distribution ratios from the equilibrium  $\frac{\text{CaO} + \text{MgO}}{\text{SiO}_2 + \text{Al}_2\text{O}_3}$  ratios are given in Fig. 2.5.

Table 2.2 (Ref.14)

Comparison of Values of f Obtained by Morris with the Calculated Values Using Data of Sherman and Chipman

Temp.	$\frac{\text{H}_2\text{S}}{\text{H}_2} \times 10^3$	C Pet	Mn Pot	S Pet	f <sub>s</sub> (Morris)	f <sub>s</sub> =f <sub>s</sub> f <sub>s</sub> f <sub>s</sub> Mn (Calculated)
1600	2.36	0.0	2.2	1.04	0.90	0.83
1600	2.40	2.2	1.8	0.54	1.77	1.47
1600	2.42	4.0	2.8	0.29	3.32	2.63
1600	2.37	Saturated	2.8	0.17	5.55	5.46
1500	2.40	Saturated	2.4	0.19	5.84*	-
1400	2.34	Saturated	1.0	0.20	6.50*	-
				•		

<sup>\*</sup> Recalculated by using equation (2.13) and Morris's data.

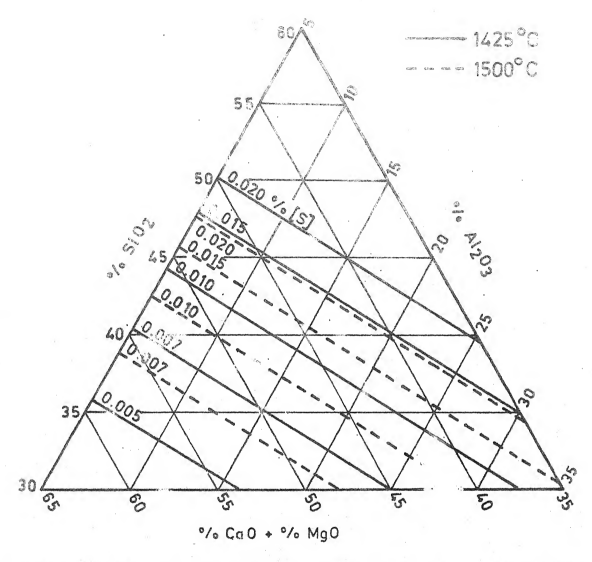


FIG. 2.4 METAL ISO-SULPHUR LINES FOR 1.5 % SULPHUR SLAGS AT 1500°C AND 1425°C. (SLAG. COMPOSITION IN WEIGHT %). (Ref. 15)

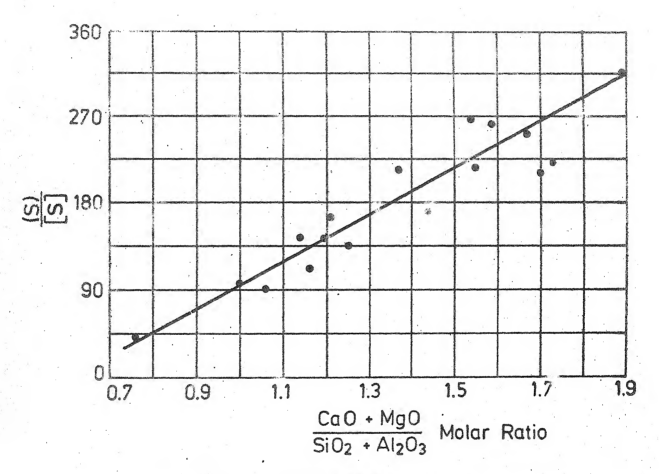


FIG. 2.5 EFFECT OF  $\frac{\text{CaO} + \text{MgO}}{\text{SiO}_2 + \text{Al}_2\text{O}_3}$  ON DESULPHURIZATION AT 1500°C. (Ref. 15)

# 2.5 Sulphur Reactions and Sulphur Control in Blast Furnace

Many studies are reported in the literature to enhance understanding of the sulphur reactions occurring in iron and steel making processes 16-25. Over all sulphur transfer between slag and metal can be written as follows:

$$[FeS] + (CaO) = (CaS) + (FeO)$$
 (2.21)

The equilibrium constant for reaction (2.21) can be expressed as follows;

$$K_{2.21} = \frac{(a_{GaS}) (a_{FeO})}{[a_{FeS}] (a_{GaO})}$$
 (2.22)

where  $a_{CaS}$ ,  $a_{FeO}$ ,  $a_{CaO}$ ,  $a_{FeS}$  represent the activity of CaS, FeO, CaO in the slag phase and that of FeS in the metal phase respectively.

FeO in slag can further combine with carbon or silicon in iron to give CO gas or SiO<sub>2</sub> in slag. The over all sulphur transfer reaction in presence of carbon can thus be written as follows<sup>26</sup>;

$$CaO(s) + S(1 Pet) + O(gr) = CaS(s) + OO(g)$$
 (2.23)

The equilibrium constant  $K_s$  for reaction (2.23) is written as follows;

$$K_{s} = \frac{a_{CaS}}{a_{CaO}[Pct. S]f_{s}} \times \frac{P_{CO}}{a_{C}}$$
 (2.24)

where  $f_s$  is the activity co-efficient of sulphur in the melt and it is believed to be around 7 for carbon saturated iron,  $a_C$  is activity of carbon in melt and may be taken as unity for

carbon saturated iron,  $p_{CO}$  is partial pressure of CO. The temperature dependence of the equilibrium constant  $K_s$  is determined by finding the standard free energy change of the reaction (2.23) by knowing the free energy of formation of each species and is given by;

$$\log K_{s} = -\frac{6010}{T} + 5.935 \tag{2.25}$$

Data on standard free energy of solution and formation of the oxides and sulphides are given in Tables 2.3 and 2.4. The activities of CaS and CaO relative to pure solids may be represented in terms of their mole fraction and activity coefficients;

$$\frac{a_{CaS}}{a_{CaO}} = \frac{56}{32} \times \frac{(Wt.Pct.S)}{(Wt.Pct.CaO)} \frac{\gamma_{CaS}}{\gamma_{CaO}}$$
(2.26)

Simplifying equation (2.24) for carbon saturated iron in a blast furnace,

$$\frac{\text{(Wt.Pct.S)}}{\text{[Wt.Pct.S]}} \quad p_{\text{CO}} = 4 \left[ \frac{\gamma_{\text{CaO}}}{\gamma_{\text{CaS}}} \right] \text{ (Wt.Pct.CaO)} \quad 10^{-\frac{6010}{\text{T}}} + 5.935$$

$$(2.27)$$

Values of  $\gamma_{\rm CaO}/\gamma_{\rm CaS}$  ratio at different  $\frac{{\rm CaO}}{{\rm SiO}_2}$  ratio of the slag are taken from the work of Sharma and Richardson and the equation (2.27) is plotted in Figure 2.6. In the bosh region of the blast furnace CO content of the gas is around 35 pct-and p<sub>CO</sub> will be very close to 1 atm. At temperature of 1500°C at the bosh, the sulphur distribution ratio,

Table 2.3

Standard Free Energy of Solution of Various
Elements in Liquid Iron

M(pure) = M (1 percent solution, Hypothetical)

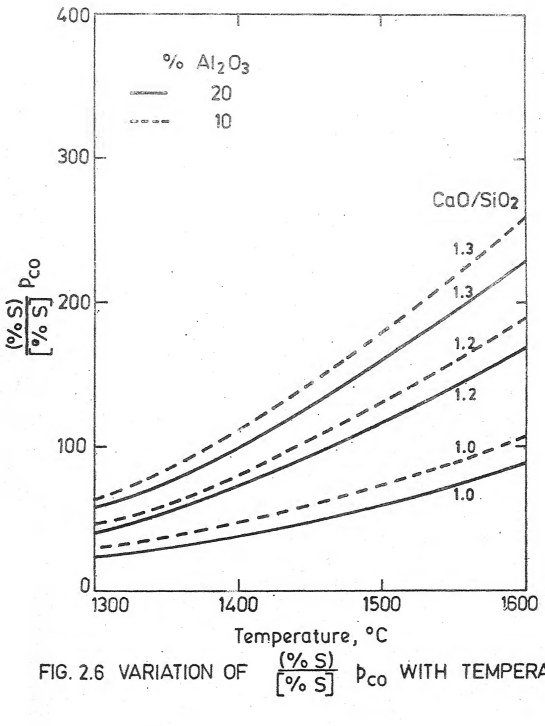
(Ref. 29)

Element, State	$\Delta$ G $^{ m o}$ Cal
C(gr) = C(Wt%)	5,100 - 10.00 T
Mn(1) = Mn(Wt%)	- 9.11 T
$\frac{1}{2} \circ_{2}(g) = 0 (Wt \%)$	-28,000 - 0.69 T
$Si(1) = \underline{S}i \text{ (Wt%)}$	-28,500 - 6.09 T
$\frac{1}{2} S_2(g) = \underline{S}(Wt\%)$	-31,520 + 5.27 T

Standard Free Energies of Formation of Compounds at Steel Making Temperatures Ref. 29.

$\nabla G_{o} =$	A	+	BT	Cal.
------------------	---	---	----	------

KCal.
1.0
1.0
1.5
3.0
1.0
0.5
1.0
- Carrier
2.0
2.0



\$ co WITH TEMPERATURE. FIG. 2.6 VARIATION OF

i.e.  $\frac{(s)}{s}$  should be above 100 for  $\frac{cao}{sio_2}$  ratio greater than

1.2, compared to actual range of 20 to 50 in modern blast furnaces. If  $p_{CO}$  is taken to be 2-4 atm. the calculated values will agree well with the actual value.  $p_{CO}$  values of 2-4 atm. may exist in the hearth of the furnace at slag metal interface where CO evolved is not diluted by  $N_2$  in the blast.

It may thus be concluded that slag — metal equilibrium is more or less reached in the hearth of blast furnace where small metal droplets trickle down a depth of molten slag layer. This equilibrium is not easily reached in the bosh. Similar arguments also explain the observed silicon and manganese levels in modern blast furnaces where slag metal equilibrium may be assumed to be prevailing in the hearth at  $p_{(i)}$  values of 3-4 atm.

Some confusion exists in the literature as silicon and manganese reduction reaction are believed to be sluggish and sulphur equilibrium is not believed to reach equilibrium. These arguments are valid if  $p_{\rm CO}$  is taken to be around 1 atm. only. Many workers therefore proposed different mechanisms

for transfer of sulphur from metal to the slag phase <sup>26</sup>, <sup>28</sup>. In the slag sulphur may be present as 5<sup>-</sup> ions, oxygen as 0<sup>-</sup> and Mn as Mn<sup>++</sup> and so on. Electrical neutrality of the slag must be maintained for any reaction to proceed. Thus King and Ramadhandran <sup>28</sup> demonstrated that transfer of sulphur from the metal to slag must be accompanied by the transfer of Si and Mn to the slag. Turkdogan <sup>26</sup> analysed large plant data and found more agreement for the sulphur and manganese coupled reaction, i.e.

$$[Mn] + [S] + (GaO) = (GaS) + (MnO)$$
 (2.28)

$$K_{2.28} = \frac{a_{\text{CaS}} a_{\text{MnO}}}{f_{\text{Mn}} f_{\text{s}}[\text{Mn}][\text{s}] a_{\text{GaO}}}$$
(2.29)

$$\log K_{2.28} = \frac{9080}{T} - 4.965$$
 (2.30)

The plot of [Mn]/ (Mn0) versus  $\frac{(S)}{[S]}$  at 1500°C shown in Figure 2.7. Reaction (2.28) will be equivalent to reaction (2.23) if [Mn] in metal is assumed to be governed by the following reaction;

$$(Mn0) + C(gr.) = [Mn] + CO(g.)$$
 (2.31)

As explained earlier p co must be taken to be 3-4 atm. in

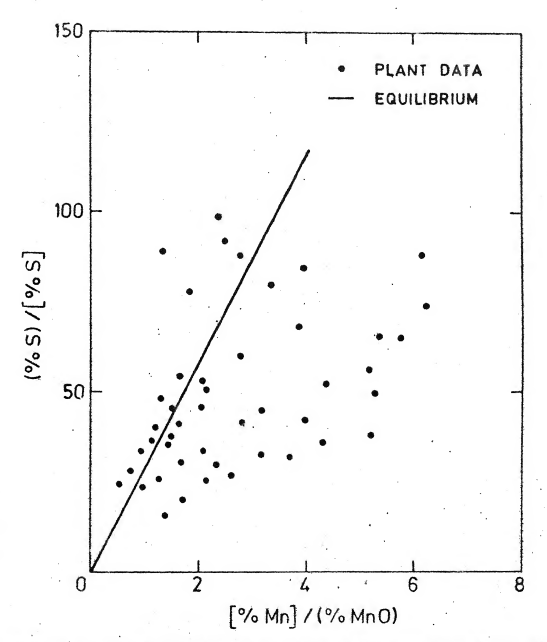


FIG. 2.7 EQUILIBRIUM SULPHUR AND MANGANESE DISTRI-BUTION RATIOS AT 1500°C COMPARED WITH PLANT DATA. (Ref. 26)

order to calculate the theoretical [Mn] content of metal for better comparison with the actual data for a blast furnace.

Similarly, the reaction involving silicon is represented by;

CaO (s) + 
$$\frac{1}{2}$$
 Si(1 Pet) +  $\frac{s}{2}$  (1 Pet) = CaS(s) +  $\frac{1}{2}$  SiO<sub>2</sub>(s)
(2.32)

The temperature dependence of the equilibrium constant  ${\rm K_{Si-S}} = \frac{{{{\left( {{a_{GaS}}} \right)}}}\left( {{a_{Si\,0}\,2}} \right)^{1/2}}{{{{\left( {{a_{GaO}}} \right)}}\left[ {{a_{Si}}} \right]^{1/2}}{{{\left( {{a_{Si}}} \right)}^{1/2}}{{\left( {{a_{Si}}} \right)}^{1/2}}} \right]} \ \, {\rm can \ \, be \ \, determined \ \, as \ \, follows,}$ 

$$\log K_{Si-S} = + \frac{9865}{T} - 4.52 \tag{2.33}$$

(S) [Si] ratio is plotted versus 
$$\frac{[Si]}{(SiO_2)}$$
 in Figure 2.8.

In some blast furnaces operated with high ash coke; S, Si, Mn reactions may not reach equilibrium with respect to carbon due to a different mechanism for the reduction of SiO<sub>2</sub> i.e. due to formation of SiO, SiS vapors at the tuyerrof a blast furnace <sup>26,36</sup>. The coupled reactions described by equations (2.28) and (2.32) may however still try to reach equilibrium in the hearth.

'Mn' content of metal will be governed by the following reaction:

$$[Si] + 2(Mn0) = 2[Mn] + (Si02)$$
 (2.34)

$$K_{Mn-Si} = \frac{(a_{SiO_2})[a_{Mn}]}{[a_{Si}](a_{MnO})^2}$$
 (2.35)

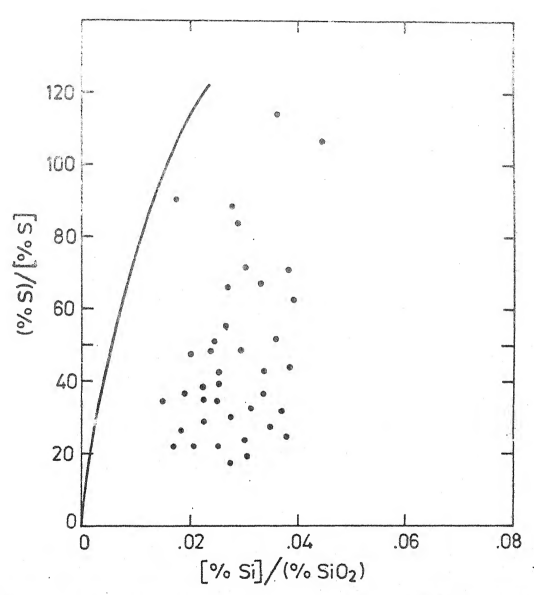


FIG. 2.8 EQUILIBRIUM SULPHUR AND SILICON DISTRIBUTION RATIOS AT 1500°C, COMPARED WITH PLANT DATA. (Ref. 26)

$$\log K_{\text{Mn-Si}} = \frac{1570}{\pi} + 0.89$$
 (2.36)

In such cases silicon replaces carbon as a reducing agent for the reduction of MnO,  $\text{Cr}_2\text{O}_3$ ,  $\text{TiO}_2$  etc.

# 2.6 Effect of Temperature on Solutbility of Sulphur in Presence of Mn

# 2.6.1 <u>Iaboratory studies</u>

Oelsen<sup>6</sup> studied the solubility of MnS in carbon saturated blast furnace iron and reported that the solubility of MnS decreased in the melt with decreasing temperature as shown in Figure 2.9. 'Si', and 'P' content of the blast furnace iron were not specified by Oelsen.

Morris 14 studied Mn-S equilibrium in Fe-C alloys by stepwise addition of manganese. His results are summ@rised in Figure 2.10. Morris reported that [Mn%][S%] product was nearly constant over a concentration range of 1 to 7 Pct. Mn, at a fixed temperature. The average values of [Mn%][S%] products were 0.05 and 0.26 at temperatures of 1200°C and 1400°C respectively, matching well with Oelsen's work. At 1600°C the [Mn%][S%] product increased from 0.8 at 1 pct Mn to 1.2 at 7 pct. Mn. Where as according to Oelsen's [Mn%][S%] product increased from 0.57 at 1 pct. Mn to 0.92 at 4 pct. Mn, at the same temperature.

Morris concluded from his data that manganese alone could not be an efficient desulphuriser except at temperatures

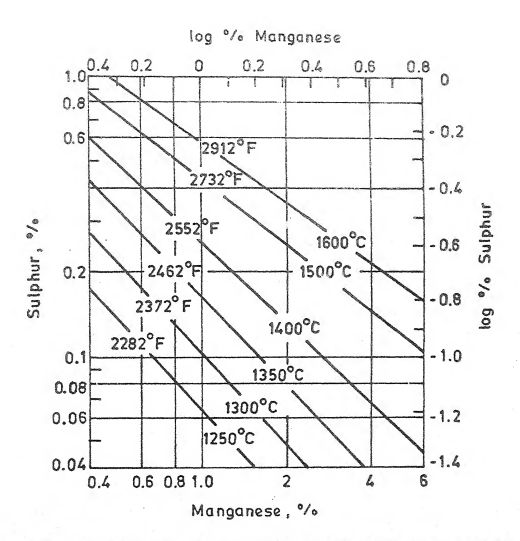


FIG. 2.9 EFFECT OF MANGANESE ON SOLUBILITY OF MANGANESE SULFIDE IN HOT METAL. (Ref. 6)

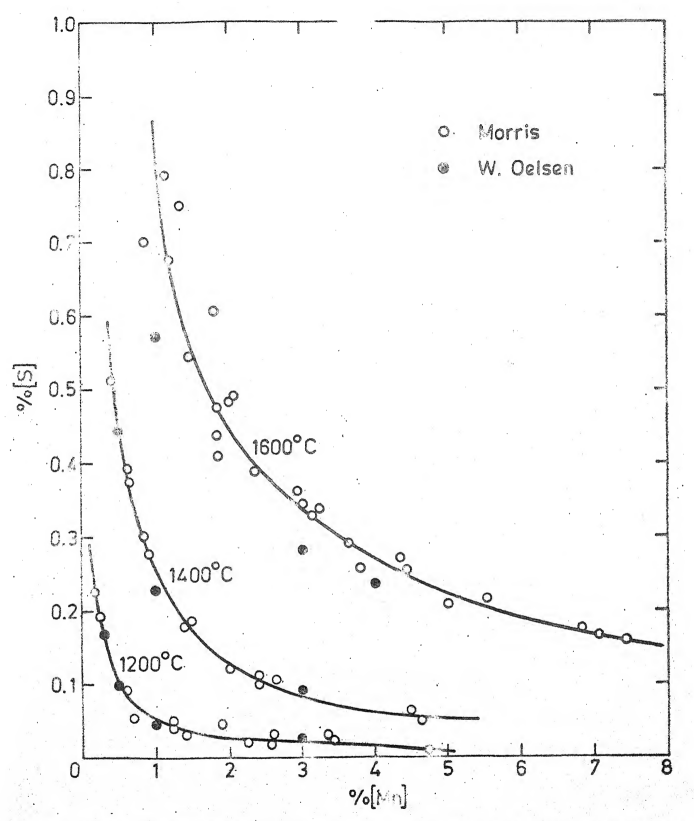


FIG. 2.10 EFFECT OF MANGANESE ON THE SULPHUR CONTENT OF CARBON SATURATED MELTS OF IRON IN EQUILIBRIUM WITH MnS-FeS SLAG. (Ref. 14)

below the normal operating range. However he predicted that in the presence of an exide slag, manganese could be more effective as a desulphuriser than indicated in his results, due to lowering of the activity of manganese sulphide in the slag. This aspect of decreasing  $a_{MnS}$  (i.e. activity of MnS) in slag was not studied by him.

#### 2.6.2 Industrial data

If initial sulphur level in iron is high, and the temperature of metal decreases during the transit of the ladle, some sulphur may precipitate out as MnS. Corresponding drop in Mn content of metal may be too small to be noted as the concentration of sulphur does not normally exceed 0.05 Pct. Drop in sulphur may be of the order of 0.01 to 0.02 Pct. and it can account for 30 to 50 Pct.of sulphur removal in the slag. Rourkela steel plant has been producing the blast furnace metal with Mn contents of 1.5 to 2.5 Pat. consistently over the last many years. There is a substantial difference in sulphur levels of the samples taken at the time at blast furnace tapping and at the time of taking out metal from the mixer. Typical Mn and S levels in blast furnace iron and mixer iron are plotted in Figure 2.11 for comparison purposes. More details regarding metal temperature, mixer slag composition etc. are not available. Slag is likely to contain CaO, SiO2, Al<sub>2</sub>0<sub>3</sub>, MnO, FeO, and sulphur as its origin may be due to the carry over slag from the blast furnace, oxidation of metal in the mixer, addition of coke breeze to the top layer of iron ladle, formation of MnS due to lowering of temperature etc.

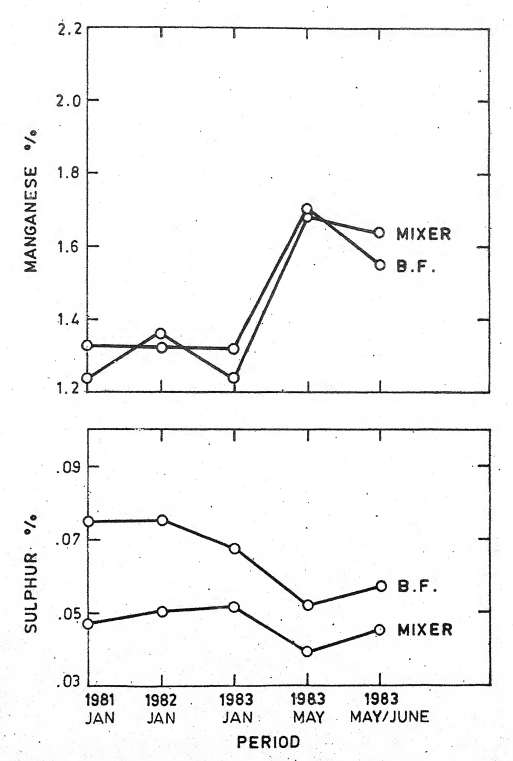


FIG. 2.11 INDUSTRIAL DATA ON MANGANESE AND SULPHUR LEVELS IN BLAST FURNACE IRON AND MIXER IRON. (SOURCE - ROURKELA STEEL PLANT)

A typical mixer slag composition is given below; CaO = 20%,  $SiO_2$ , 40%,  $Al_2O_3 = 20\%$ 

FeO = 5%, S = 0.70% and the balance may be Fe, MnO, MgO etc. This slag must be removed carefully as to take full advantage of sulphur removal as MnS, otherwise sulphur would revert to metal under oxidising condition in the LD steel making where metal temperature may rise to  $1600^{\circ}$ C. Rourkela Steel Plant has acquired 3 slag racking machines for this purpose.

Mn may not act as desulphuriser alone to attain the lower 'S' levels in metal and it may be necessary to add desulphurising agents such as CaO, CaO2, Na2CO3 etc.

Rourkela steel plant has incorporated the facility of external desulphurisation where CaO2 powder is injected to metal using coated lances and air as carrier. Other plants like Bhilai, Bokaro may also have such facilities in their expansion programmes. It is not known how the presence of Mn in metal will influence the desulphurisation capacity of these agents. Results of the laboratory investigations reported in the present work may be helpful in better understanding of the role of Mn in the external desulphurisation of iron.

#### CHAPTER III

#### PLAN FOR PRESENT WORK

From the available literature and the industrial data, practical importance of 'Mn' and 'S' reactions in blast furnace iron can not be under estimated. However, few scientific studies are available in the temperature range of 1200°C to 1400°C, which may usually occur in the transit of metal from the blast furnace to the mixer and during external desulphurisation of iron. The present work was aimed at making such studies under controlled conditions in the laboratory. The work may be described as follows;

- (1) The literature survey comprising the critical evaluation of the necessary thermodynamic data.
- (2) Laboratory experiments to establish relationships between 'Mn' and 'S' in the Fe-C alloys in temperature range of 1200° 1400° C with or without any slag additions. This included fabricating a silicon carbide furnace, making Fe-C alloys, melting the alloy under argon atmosphere in a graphite crucible, drawing samples with a quartz tube and careful analysis of sulphur and manganese in the laboratory.
- (3) The effect of 'Mn' in metal on the desulphurising power of the oxide slags was studied by making synthetic slag of CaO-SiO<sub>2</sub>-Al<sub>2</sub>O<sub>3</sub> in the laboratory and equilibrating the melt at 1400°C with those slags for 4-6 hours of duration. Samples were either drawn from the

melt or taken from the solidified mass in the crucible after the duration of experiment. Pure lime was also used as a desulphuriser in some experiments for comparison purposes.

- (4) Evaluation of the laboratory data by developing suitable theoretical and multicomponent models and by comparison with the other published data in such system.
- (5) Collection and evaluation of industrial data to signify the practical importance of the present work.

#### CHAPTER IV

#### RESULTS OF THEORETICAL CALCULATIONS

# 4.1 Evaluation of Reported Thermodynamic Data

\*As mentioned in Section 2.6.1, solubility of sulphur in iron carbon alloys decreases with addition of Mn to the alloy at a constant temperature and with decrease in temperature at a constant Mn level. It is also possible to determine theoretically the sulphur content of metal as function of temperature and Mn content of metal if the corresponding data on standard free energy of formation of MnS and the activity coefficient of 'Mn' and 'S' in metal are known.

# 4.1.1 Standard free energy of formation

The standard free energy change for the following reaction can be determined from the values given in Tables 2.3 and 2.4.

$$\underline{\mathbf{Mn}} (\mathbf{Wt\%}) + \underline{\mathbf{S}} (\mathbf{Wt\%}) = \mathbf{MnS(1)}$$
 (4.1)

$$\Delta G_{4.1}^{0} = -31,580 + 19.61 \text{ T Cal.}$$
 (4.2)

 $\Delta G_{4.1}^{0}$  is valid in the temperature range  $1803^{0}-2000^{0}$ K The equilibrium constant  $K_{4.1}$  for the reaction (4.1) is given by;

$$K_{4.1} = \frac{a_{MnS}}{f_{Mn} f_{s} [Wt\% Mn][Wt\% S]}$$
 (4.3)

and 
$$\log \frac{a_{MnS}}{f_{Mn} f_{s} [wt\% Mn] [wt\% S]} = \frac{6902.7}{T} - 4.28$$
 (4.4)

In the temperature range of 15160 - 18030K,

$$\underline{Mn}(wt) + \underline{S}(wt\%) = \underline{MnS}(s)$$
 (4.5)

and 
$$\triangle G_{4.5}^{0} = -37,730 + 23.02 \text{ T cal}$$
 (4.6)

where,  $a_{MnS}$  is the activity of MnS is slag,  $f_{Mn}$  and  $f_s$  are the activity coefficients of [Mn] and [S] in the melt. The standard states of mangane and sulphur are chosen to be 1 wt pct in the solution and their activity coefficients approach unity when the concentrations of all the solutes tend to be zero.

# 4.1.2 Activity coefficient of sulphur

For finite concentrations of solutes, one has to know the value of interaction parameters to determine the activity coefficient as follows;

$$\log f_{S} = \sum_{i=1}^{n} e_{S}^{i} (wt\% 1)$$
 (4.7)

Values of interaction parameters have been determined by various workers  $^{7,10}$ , by equilibrating the melt with  $\rm H_2$  and  $\rm H_2S$  gas mixtures and determining the sulphur content of the metal. Table 4.1 summarises the interaction parameters at  $1600^{\circ}\rm C$ .

Table 4.1

Values of Interaction Parameters at 1600°C

		•		
Solute j	e <sup>j</sup> s	Ref.	e j Mn	Ref.
	0.11	30	and the state of t	
	0.12	12		
C	0.15	10	-0.07	30
	0.24	31		
Chief Cuth Little Common and Chief State Carpet Source Chief State	0.063	30	THE CONTROL COURSE STATE SECURITY PROPERTY AND ADMINISTRATION OF THE COURSE STATE STATE OF THE COURSE STAT	
Si	0.066	31	0.0	30
Mn	-0.026	30	0.0	32 30
armen. Sipt 4-little verkjone godine mengript grave visigne	0.045	31		Targeter (produces of the AFE). The East of Target Eastern Age of Eastern Age of Eastern Eastern Eastern Easte
P	0.029	30	-0.0035	30
S	-0.028	30,31	-0.048	30
			-0.043	31

For carbon saturation, pct carbon is very high compared to other solutes and because of this carbon-sulphur interactions are important,  $e_s^c$  values should be thus known more accurately than other values. Unfortunately there is scatter in the data shown in Table 4.1. Sources of discrepancies in  $e_s^c$  values have already been discussed in Chapter 2.

Young and Clark  $^{33}$  recently established that  $e_s^c$  value of 0.16  $\pm$  0.01 at  $1600^o$ C gives the best approximation to the actual plant data. This is closer to the average value of 0.15 from the data of Morris and Buehl  $^{10}$ .

# 4.1.3 Activity coefficient of manganese

It is generally assumed that most alloying elements have no significant effect on the ideal behaviour of manganese in steel, but carbon is known to decrease the activity of manganese in iron-carbon-manganese alloys  $^{34,35}$ . The variation of activity coefficient  $f_{\rm Mn}$  with composition in liquid Fe-C-Mn alloys at  $1540^{\circ}$ C is shown in Figure 4.1.

From Figure 4.1 it is seen that for carbon saturated iron containing 3 pct Mn (broken line  $X_{\rm Mn}$ =0.03),  $f_{\rm Mn}^{\rm c}$  value is around 0.95. Turkdogan has estimated the activity coefficient of Mn in carbon saturated liquid iron to be 0.8.

In the present work  $f_{Mn}$  is assumed to be 0.8, for the carbon saturated melt in the absence of more accurate data.

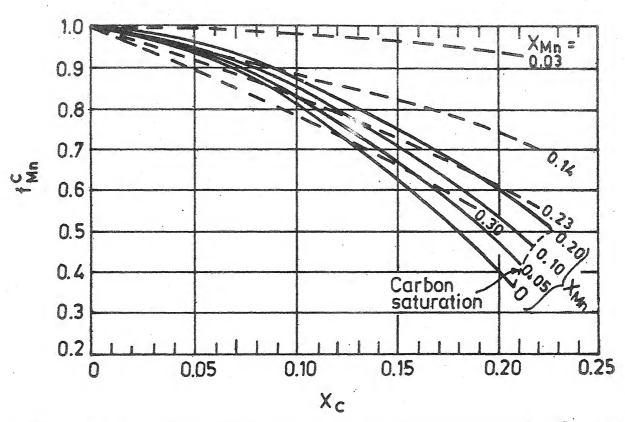


FIG. 4.1 VARIATION OF THE ACTIVITY COEFFICIENT, I No., WITH COMPOSITION IN LIQUID IRON-CARBON-MANGANESE ALLOY AT 1450°C. (Ref. 34, 35)

(FULL CURVE Ref. 35 AND BROKEN CURVE Ref. 34)

# 4.2 Determination of Activity Coefficients of Sulphur and Manganese at Temperatures of 1200°C to 1500°C

Melting point of iron decreases in presence of carbon. For carbon saturated iron, melting point may be as low as 1150°C. Most of the thermodynamic data is however obtained at 1600°C with reference to the steel making conditions. It is necessary to know the values of activity coefficients at lower temperatures of 1200° - 1500°C if the solubility of sulphur is to be calculated in iron-carbon alloys at temperatures less than 1600°C. In the present work, a regular solution model is applied to extrapolate the interaction parameters.

The expression for the standard free energies of formation given in Table 2.3 - 3.4 are assumed to be valid at these temperatures though they might have been derived from the experimental data in the temperature range 1500°-1600°C.

For a regular solution the quantity RT ln 7; should not change with temperature, where R is the gas constant, T is temperature in K and  $\gamma$  is the activity coefficient of the solute 'i' in the Raoultian scale.

The Rapultian and Henrian interaction coefficients are defined as follows:

$$e_{\mathbf{i}}^{\mathbf{j}} = \frac{\partial (\ln \gamma_{\mathbf{i}})}{\partial (\mathbf{x}_{\mathbf{j}})} \quad \mathbf{x}_{\mathbf{i}}, \mathbf{x}_{\mathbf{j}} \to 0$$

$$e_{\mathbf{i}}^{\mathbf{j}} = \frac{\partial (\log f_{\mathbf{i}})}{\partial (\operatorname{wt}_{\%} \mathbf{j})} \quad \operatorname{wt}_{\%} \mathbf{i}, \mathbf{j} \to 0$$
(4.8)

$$\mathbf{e_i^j} = \frac{\partial(\log f_i)}{\partial(\text{wt}\% j)} \text{ wt}\% i, j \to 0$$
 (4.9)

The term  $e_i^j$  corresponds to the reference state of the pure substance, and concentrations are expressed in atom or mole fractions. The term  $e_i^j$  is applicable to the infinitesimal dilute solution as the reference state and concentrations are expressed in weight percents.

As RT  $\ln \gamma_i$  is constant for a regular solution ; one can write the following;

$$RT_1 \in {}_{\mathbf{i}}^{\mathbf{j}}(1) = RT_2 \in {}_{\mathbf{i}}^{\mathbf{j}}(2)$$
 (4.10)

where  $\epsilon_{i}^{j}(1)$  and  $\epsilon_{i}^{j}(2)$  are Raoultian interaction parameters at temperatures  $T_{1}$  and  $T_{2}$  respectively. It is also known that, the two types of interaction parameters are related as follows<sup>31</sup>;

$$\hat{\mathbf{e}}_{\mathbf{i}}^{\mathbf{j}} = 230 \, \frac{M_{\mathbf{j}}}{M_{\mathbf{1}}} - \hat{\mathbf{e}}_{\mathbf{i}}^{\mathbf{j}} + \frac{M_{\mathbf{1}} - M_{\mathbf{j}}}{M_{\mathbf{1}}}$$
(4.11)

 $M_j$ ,  $M_l$  are the molecular weights of the solute  $l_j$  and the solvent iron.

Let 
$$\frac{M_1 - M_j}{M_1} = C_0$$
 (4.12)

On simplification equation (4.11) can be written as;

$$\frac{1 - \frac{C_0}{e_{\mathbf{i}}^{\mathbf{j}}(1)}}{\frac{T_0}{\mathbf{i}}} = \frac{T_1}{T_2} \frac{e_{\mathbf{i}}^{\mathbf{j}}(1)}{e_{\mathbf{i}}^{\mathbf{j}}(2)}$$

$$1 - \left[\frac{T_2}{T_1} \times \frac{C_0}{e_{\mathbf{i}}^{\mathbf{j}}(1)}\right]$$
(4.13)

Henrian interaction coefficient  $e_{\mathbf{i}}^{\mathbf{j}}(1)$  at temperature  $T_{\mathbf{i}}$  (i.e.  $1873^{\mathbf{o}}$ K) is known from literature (Table 4.1). So at any other temperature  $T_{\mathbf{i}}$ ;  $e_{\mathbf{i}}^{\mathbf{j}}(2)$  i.e. the interaction coefficient at the desired temperature can be calculated by using equation (4.13).

The results of the calculations of interaction parameters are summarised in Table 4.2. Using these values  $f_s$  can be determined at any temperature for any batch composition with the help of equation 4.7 and  $f_{Mn}$  can be determined using the following.

$$\log f_{Mn} = \Sigma e_{Mn}^{i} (wt\% i)$$
 (4.14)

value of  $f_{Mn}$  is, however, assumed to be 0.80 for carbon saturated iron at all temperatures.

# 4.3 Solubility of Sulphur in Fe-Mn-S System at 1600°C

# 4.3.1 Determination of $f_s$ and $f_{Mn}$ values

The  $f_s$  and  $f_{Mn}$  values are determined by taking the interaction parameter values as given in Table 4.1 and applying equation (4.7) and (4.14).

# 4.3.2 Determination of equilibrium sulphur levels

Equilibrium sulphur level of the metal at 1600°C is calculated by using equation (4.4). The procedure of calculation is discussed below. In the first case the slag is assumed to consist of only manganese sulphide so that a<sub>MnS</sub> can be taken as unity.

For a particular Mn level in the batch S level is first assumed or estimated approximately  $\mathbf{f_s}$  and  $\mathbf{f_{Mn}}$  values are then calculated by using equation (4.7) and equation (4.14); and the sulphur level is recalculated using equation (4.4).

Table 4.2  $e_i^j$  Values at  $1200^{\circ}\text{C} - 1600^{\circ}\text{C}$ 

Temp.	e c s	e <sup>Mn</sup> s	e <sup>s</sup> s
1600	+ .1500	0260	-,0280
1500	+ .1600	0270	0295
1400	+ .1698	0290	0309
1300	+ .1816	0309	0327
1200	+ .1950	0330	0347

The steps are repeated until the successive calculated sulphur levels are about the same.

In the second case the slag is assumed to consist of iron sulphide and manganese sulphide binary system and their behaviour is assumed to be ideal at 1600°C. From the FeS-MnS phase diagram as shown is Figure 4.2, it is seen that FeS and MnS are completely soluble in the liquid phase. Activity of FeS or its mole fraction in the slag is related to sulphur content of iron as follows;

$$\underline{F}_{0} + \underline{S} = \underline{F}_{0}\underline{S} \tag{4.15}$$

$$K_{4.15} = \frac{a_{FeS}}{a_{Fe} f_{s} [wt\% S]}$$
 (4.16)

Setting are equal to unity

$$a_{\text{FeS}} = K_{4.15} f_{\text{s}} [\text{wt} \% \text{ S}]$$
 (4.17)

Assuming ideal behavior, a<sub>FeS</sub> = X<sub>FeS</sub>.

For the slag consisting of FoS and MnS only

$$a_{MnS} = X_{MnS} = 1 - X_{FeS}$$
 (4.18)

where  $X_{FeS}$  and  $X_{MnS}$  are the mole fractions of FeS and MnS respectively. But from equation (4.3);

[wt % S] = 
$$\frac{e_{MnS}}{f_{Mn} f_{s} [Mn] K_{4-1}}$$
 (4.19)

By inserting equations (4.17) and (4.18) in equation (4.19) and simplifying, the sulphur level in the metal can be calculated as follows;

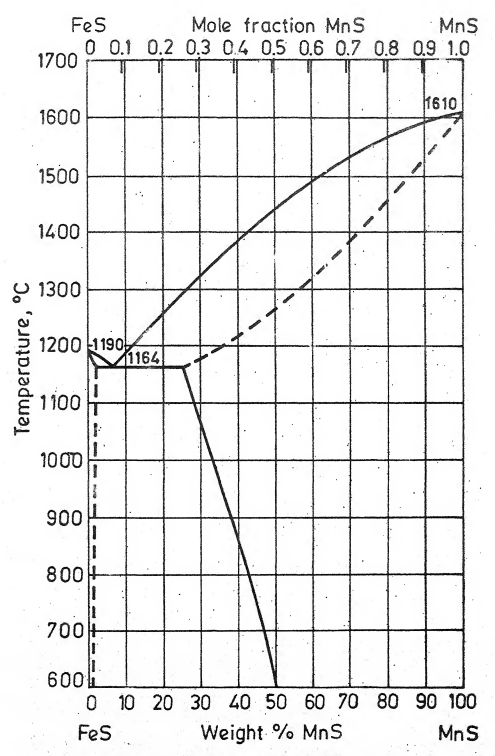


FIG. 4.2 PHASE DIAGRAM OF FeS - MnS. (Ref. 37, 38)

[wt % S] = 
$$\frac{1}{f_s(K_{4.15} + f_{Mn}[Mn]K_{4.1})}$$
 (4.20)

The values of  $f_s$ ,  $f_{Mn}$  and equilibrium sulphur content are summarised in Table 4.3.

#### 4.3.3 Comparison with published data

The calculated sulphur and manganese levels in

Table 4.3 are plotted in Figure 4.3 along with the
experimental data of Sherman and Chipman. The calculated
sulphur levels lie much above the experimental points of
Sherman and Chipman, who determined sulphur and manganese
levels in the melts after allowing them to reach equilibrium
with respect to H2+H2S gas mixtures. It is obvious that
MnS could not be formed in their experiments as sulphur
content was always below the level at which MnS or FeS would
precipitate out. Meyer s<sup>39</sup> experimental results of the
equilibrium sulphur and manganese levels in the FeS+Mn = Fe+MnS
system are also plotted in Figure 4.3. Sulphur levels in
Meyer's experiment are found to be lower than the equilibrium
values because of the presence of 4 to 6 pct. Al<sub>2</sub>O<sub>3</sub> which
might have lowered the values of a<sub>MnS</sub> and a<sub>FeS</sub> further.

# 4.4 Solubility of Sulphur in Fe-Mn-C-S- System

# 4.4.1 Solubility of carbon in iron alloys

For iron-carbon alloys it is possible to work at temperatures less than 1600°C. At any temperature saturated carbon level in the alloy can be calculated by using the following relationship<sup>33</sup>:

Table 4.3

Equilibrium[Mn]and[S]levels at 1600°C in Fe-Mn-S System

Pet[Mn]	$\mathtt{f}_{\mathtt{Mn}}$	fs	Pet[S] a <sub>MnS</sub> =1	Pct [S] a <sub>MnS</sub> =X <sub>MnS</sub>	$\mathbf{x}_{ ext{MnS}}$
2.5	0.70	0.70	3.20	2.26	0.70
3.0	0.75	0.70	2.50	1.88	0.75
4.0	0.81	0.70	1.80	1.41	0.78
5.0	0.85	0.67	1.38	1.18	0.85

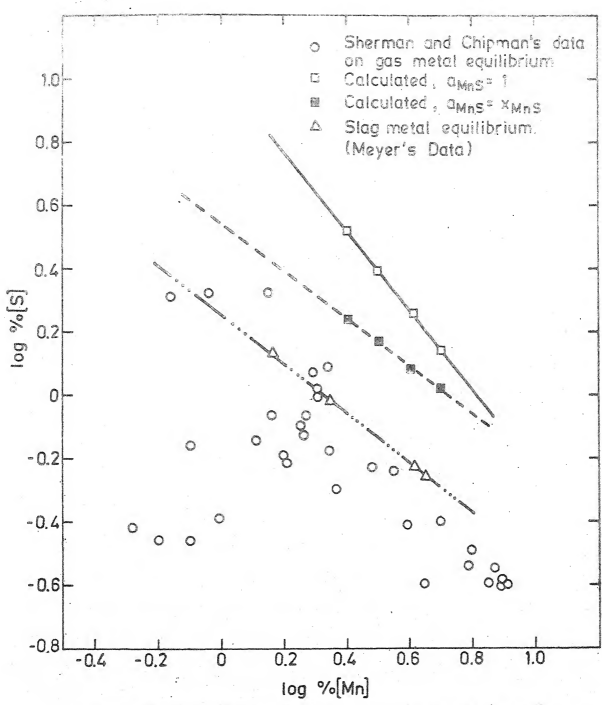


FIG. 4.3 EQUILIBRIUM [Mn] AND [S] LEVELS AT 1600°C.

$$[c]_{sat} = 1.34 + 2.54 \times 10^{-3} (T^{\circ}c)$$

$$-0.35[P] + 0.17 [Ti] - 0.40 [S]$$

$$+ 0.04[Mn] - 0.30[Si]$$
 (4.21)

where, the elements in the bracket represent their concentration in the iron alloy in wt pct.

Equation (4.21) is plotted in Figure 4.4. This shows the effect of alloying elements on carbon saturation values at different temperatures.

# 4.4.2 Determination $f_s$ and $f_{Mn}$ values

'f<sub>s</sub>' values can be determined at 1600°C by using equation (4.7) and the interaction parameter values reported in Section 4.1.2. At lower temperatures, interaction parameters can be calculated using equation (4.13), and the equation (4.7) can be then used to calculate the f<sub>s</sub> value.

For carbon saturated melt  $f_{\mbox{Mn}}$  is assumed to be 0.8 at all temperatures.

## 4.4.3 Determination of equilibrium sulphur levels

Equation (4.4) can be used to determine the sulphur level at any manganese in the metal above  $1530^{\circ}$ C, whereas equation (4.6) can be used at temperatures lower than  $1530^{\circ}$ C, provided  $f_{\rm S}$  and  $f_{\rm Mn}$  are calculated by using the suitable equations as discussed earlier. The results of the calculations by taking  $a_{\rm MnS}$  equal to unity are given in Table 4.4.

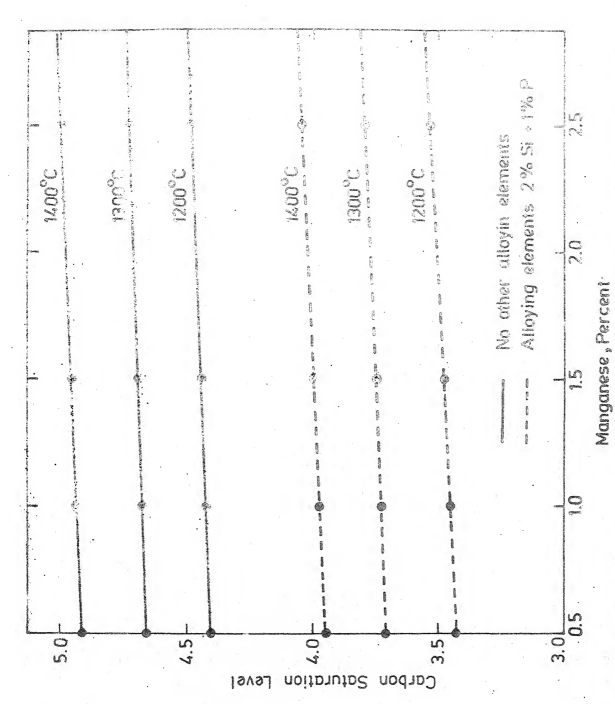


FIG. 4.4 EFFECT OF ALLOYING ELEMENTS ON CARBON SATURATION LEVEL. AT DIFFERENT TEMPERATURES.

Table 4.4 Results of Theoretical Calculations of Equilibrium [Mn] and [S] Levels in Fe-C-Mn-S System at 12000-1500°C (Assumption:  $f_{Mn}$  = 0.8 and  $a_{MnS}$  = 1.0)

Temp.	Pct[Mn]	Pct C <sub>sat</sub>	fs	Pet [S] a MnS=1	Pct[S] a <sub>MnS</sub> =X <sub>MnS</sub>
	0.4	4.38	7.05	0.120	0.1080
	1.0	4.41	7.00	0.048	0.0460
	1.4	4.43	6.98	0.034	-
1.000	2.0	4.46	6.93	0.024	0.0227
1 200	2.6	4.48	6.87	0.019	-
	3.0	4.51	6.83	0.017	0.0168
	3.6	4.53	6.78	0.014	-
	4.0	4.55	6.74	0.013	0.012
# NE CUMALANCE STATE LANCANISCISSIONE	0.4	4.60	6.86	0.280	0.2190
	1.0	4.66	6.80	0.112	0.1010
	1.4	4.67	6.76	0.080	-
1700	2.0	4.70	6.71	0.056	0.0540
1300	2.6	4.74	6.66	0.044	
	3.0	4.76	6.63	0.038	0.0370
	3.6	4:79	6.58	0.032	-
	4.0	4.82	6.55	0.029	0.0280

Contd....

Table 4.4 contd...

Temp °C	Pet Mn	Pct C <sub>Sat</sub>	fs	Pet [S] a <sub>MnS</sub> =1-	Pct[S] a <sub>MnS</sub> =X <sub>MnS</sub>
	0.4	4.89	6,68	0.587	0.364
	1.0	4.91	6,63	0.237	0.190
	1.4	4.93	6.59	0.171	-
	2.0	4.95	6.54	0.120	0.107
1400	2.6	4.97	6.48	0.093	-
	3.0	5.00	6.45	0.082	0.075
	3.6	5.02	6.40	0.068	_
	4.0	5.05	6.36	0.062	0.058
	0.4	5.14	6.52	1.130	0.500
	1.0	5.16	6.47	0.462	0.309
	1.4	5.18	6.44	0.332	_
1500	2.0	5.20	6.39	0.233	0.189
1500	2.6	5.22	6.34	0.181	_
	3.0	5.24	6.31	0.157	0.137
	3.6	5.27	6.26	0.132	- ,,
	4.0	5.30	6, 23	0.120	0.108

#### 4.4.4 Comparison of published data

f<sub>s</sub> values obtained by regular solution model in Table 4.4 agree well with f<sub>s</sub> values obtained from data of Morris <sup>14</sup> in Table 2.2. The calculated value of f<sub>s</sub> around 6.85 in Table 4.4 for 1 pct Mn in the melt at 1400°C match well with the value of 6.5 from the data of Morris <sup>14</sup>. At 1500°C for 2.4 pct Mn in the melt, the calculated value of around 6.35 is slightly greater than the experimental value of 5.64.

# 4.5 Solubility of Sulphur in Fe-Mn-C-Si-S and Fe-Mn-C-P-S-Systems

The activity coefficient of sulphur increases slightly in presence of silicon and phosphorous due to the positive values of their interaction parameters and this compensates to some extent the decrease in the activity coefficient of sulphur caused by the decrease in the solubility of carbon in the melt. The results of the calculations are given in Table 4.5.

Table 4.5

Results of Calculations of Activity Coefficients and Equilibrium Sulphur Content at Different Compositions of Melt and Temperatures

		San					
Temp og	[% Si]	[%P]	[% Mn]	[c <sub>sat</sub> ]	$\mathtt{f}_{\mathtt{Mn}}$	fs	[ %S ]
	0	0	0.5	4.9	0,8	6.63	0.476
	2	0	0.5	4.3	0.8	7.30	0.436
1.100	2	1	0.5	4.0	0.8	6.30	0.470
1400	0	0	2.5	4.9	0.8	6.00	0.105
	2	0	2.5	4.3	0.8	6.53	0.096
	2	1	2.5	4.0	0,8	6.12	0.102
ignerabiones and confidential	0	0	0.5	4.50	0.8	6.80	0.100
	2	0	0.5	<b>3.</b> 88	0.8	7.50	0.091
1200	2	1	0.5	3.50	0.8	7.00	0.097
1200	0	0	2.5	4.50	0.8	6.08	0.022
	2	0	2.5	3.88	0.8	6.73	0.020
	2	1	2.5	3.50	0.8	6.23	0.022

#### CHAPTER V

#### EXPERIMENTAL DETAILS

#### 5.1 Equipments

#### 5.1.1 Silicon carbide furnace

A silicon carbide furnace 40 cm 0.D. and 50 cm long was fabricated in the laboratory. The major parts of the furnace are schematically shown in Figure 5.1.

The inner part of the furnace is insulated by 10 cm thick hot face insulation bricks in a cylindrical fashion. Provisions are made to accommodate the furnace tube as well as silicon carbide rods inside the furnace. The outer layer of insulation consists of 5 cm thick, 85 Pct MgO, 15 Pct asbestos powder mix. This is surrounded by a metal shell which in turn is covered by a layer of asbestos cloth to reduce heat losses.

The bottom and top of the furnace are suitably insulated using 4 mm thick asbestos sheets.

The furnace tube consisted of an alumina tube 5 cm i.d. and 60 cm long with one end closed. The closed end of the alumina tube rested on a refractory base at the bottom and all the openingsat top and bottom were carefully blocked by refractory pieces and cement to minimise heat losses. Alumina chips and powder were used to fill up the alumina tube up to the high temperature zone of the furnace.

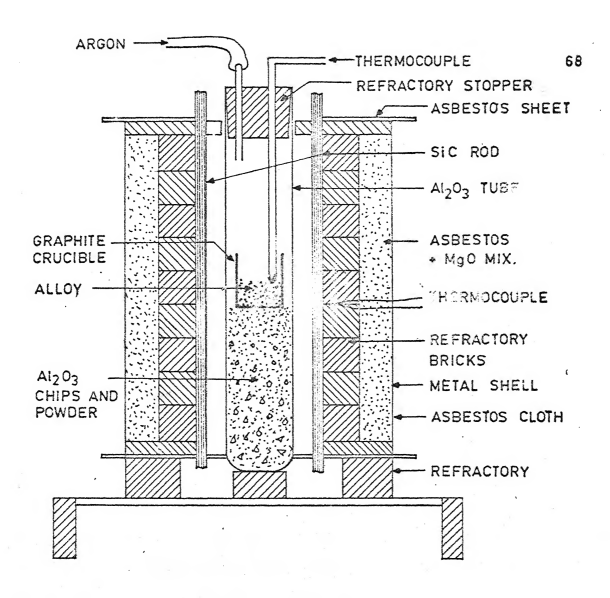


FIG. 5.1 SCHEMATIC DIAGRAM OF SILICON CARBIDE FURNACE.

#### 5.1.2 Transformer

Six silicon carbide rods, 1.8 cm dia, 55 cm long with 25 cm of hot zone, are connected in series by means of metallic connecting straps.

The power to the furnace is supplied by a continuously adjustable single phase, oil cooled, 40 A rating auto transformer with 220 volts input and 0-240 volts out put. The transformer was manufactured by 'Automatic Electric Private Ltd. Bombay'.

#### 5.1.3 Temperature controller

An ON-OFF temperature controller supplied by Leeds and Northrup Co., USA was used to regulate temperature of the furnace. A 2 ohms resistance silicon carbide furnace was connected in parallel to the relay in order to supply power to the furnace at a reduced rating when the controller was in off position. The circuit diagram is schematically shown in Figure 5.2. These arrangements helped to control the temperature of the reaction tube in the furnace within the limit of  $\pm 2^{\circ}$ C.

#### 5.1.4 Thermocouple

Platinum-Platinum 10 pct Rhodium thermocouple of 0.5 mm diameter, and 30 cm long was used to control the furnace temperature through the temperature controller. A separate 75 cm long Pt-Pt + 10% Rh. thermocouple of same diameter was used to measure the exact temperature of the melt. These thermocouples were periodically calibrated with melting point of pure copper and with a standard Pt-Pt + 10% Rh thermocouple, supplied by NPL, Delhi.

The thermocouple wires were protected by inserting them in twin hole thermocouple sheaths of appropriate length and 5 mm 0.D. The sheaths containing the thermocouple were further inserted in a 8 mm i.d. one end closed refractory tube. The outer ends of the bigger thermocouple were connected

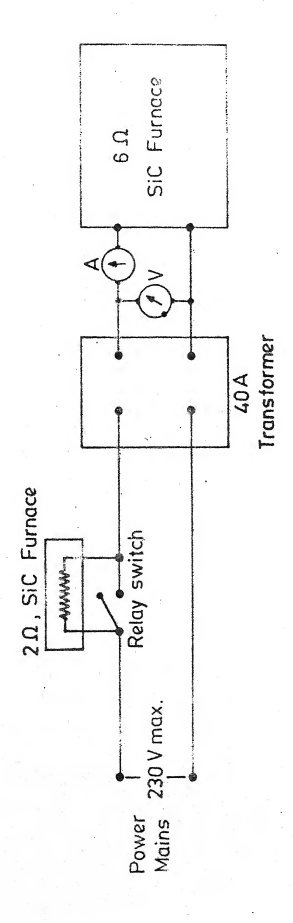


FIG. 5.2 SCHEMATIC CIRCUIT DIAGRAM FOR THE POWER SUPPLY TO THE SILICON CARBIDE FURNACE.

to the cold junction by means of compensating wires. The cold junction consisted of 5 mm i.d. glass tubes containing mercury and it was kept inside the thermos.

A thermometer with - 10° to 100°C range was used to record its temperature. The cold junction in turn was connected to the potentiometer terminals by means of copper wires, for measuring the emf generated by the thermocouple.

#### 5.1.5 Potentiometer

The millivolt potentiometer manufactured by Leeds and Northrup, Co. U.S.A. was used to measure the emf generated by the thermocouples and this in turn could be converted to temperature by using standard tables.

#### 5.1.6 Gas train

The neutral atmosphere of argon was maintained in most of experiments to protect the graphite crucible and the melt from the atmospheric oxidation. The argon gas was purified by passing through a calcium chloride tower to remove moisture and a copper gauze tube, maintained at  $260^{\circ}-300^{\circ}$ C inside a furnace to absorb oxygen.

A calibrated capillary flow meter was used to determine the flow rate of argon being passed to the furnace through the gas train.

#### 5.1.7 Induction furnace

The vacuum induction furnace having a capacity to melt around 500 gm of alloy, available in the crystal growth laboratory of IIT Kanpur, manufactured by Ajax, USA was used

to make the Fe-C alloys in graphite crucibles.

#### 5.1.8 X-ray diffractometer

The X-ray diffractometer manufactured by Rich Seifert and Co., W. Germany, stationed in the ACMS Laboratory, IIT Kanpur was used for characterisation of input and output materials.

# 5.1.9 Atomic absorption spectro-photometer (AAS)

Model No. IL 751, AAS manufactured by 'Instrumentation Laboratory, U.S.A., was used to analyse manganese in the sample.

#### 5.1.10 Sulphur analysing setup

For analysing sulphur in the iron samples, the setup was made available near the experimental site. It consisted of 250 cc conical flask, B/34 glass joint, 250 cc separating funnel, glass tubes (6 mm i.d.), 100 cc measuring cylinder, and 1500 watt hot plate. The volumetric parts for titration had 50 cc burrette and 10 cc pipette. Schematic diagram of the set up is shown in Fig. 5.3.

#### 5.2 Materials

#### 5.2.1 Steel samples

Steel samples of 5 cm diameter and 3 cm long were collected from the open hearth bath of Durgapur Steel plant, at the end of refining, but just before addition of any ferro alloys etc.

The analyses of these samples as supplied by Durgapur Steel Plant are given in Table 5.1.

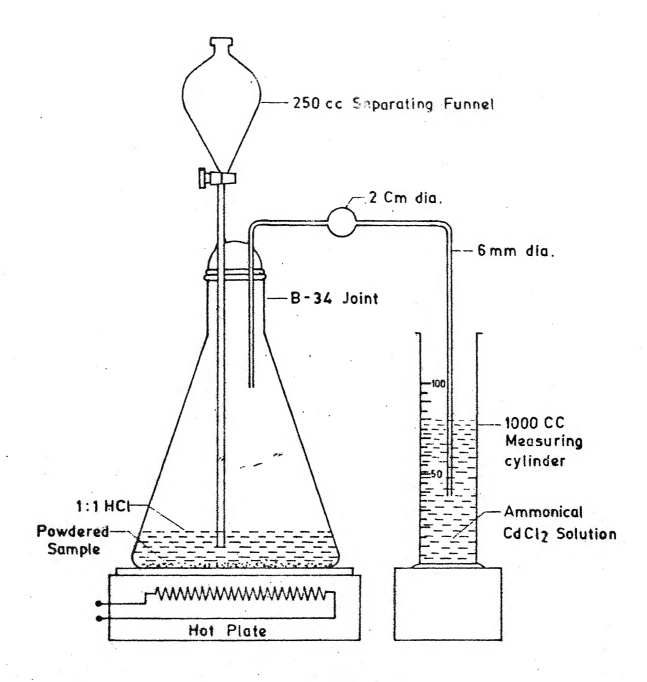


FIG. 5.3 SCHEMATIC SET UP FOR SULPHUR ANALYSIS.

Table 5.1

Chemical Analysis of Steel Samples Obtained from Durgapur Steel Plant

		Elem	ents, po	et.	
Sample No.	С	S	P	Mn	Si
1	0.12	0.048	0.020	0.05	< 0.05 for
2	0.11	0.046	0.015	for all samples	all samples
3	0.17	0.040	0.037		
4	0.14	0.038	0.015		
5	0.07	0.040	0.010		
6	0.14	0.037	0.010		
7	0.09	0.042	0.010		
8 ,	0.20	0.040	0.025		
9	0.14	0.040	0.024		
10	0.14	0.037	0.015		
11	0.14	0.042	0.016		
12	0.12	0.036	0.020		
13	0.12	0.044	0.028		
14	0.09	0.040	0.035		
15	0.13	0.040	0.010		

#### 5.2.2 Mn.FeS.MnS

99.9 Pct pure FeS and Mn (imported)were used for the necessary addition to the melt. In some cases FeS and MnS were obtained locally and added to melt. All raw materials were characterised using X-ray diffractometer.

# 5.2.3 $GaO, SiO_2, Al_2O_3$

99.0 Pct pure CaO, SiO<sub>2</sub>, Al<sub>2</sub>O<sub>3</sub> were used for making slags. Time was obtained by calcination of calcite material in the laboratory at 1000 °C for one hour.

#### 5.2.4 Graphite crucible

Graphite crucibles of 2.5 cm i.d., 7.5 cm long and 4.0 cm i.d., 10 cm long were made from 4 and 5 cm diameter locally available graphite rods, respectively.

Some crucibles were made from the broken graphite electrodes supplied by J.K. Iron and Steel, Kanpur and Singh Engineering Works, Kanpur. 5 mm holes were made at the top, sides of the crucibles to facilitate lowering of the crucible to the furnace tube using the hook of a long steel rod.

#### 5.2.5 Sampling tube

4 to 6 mm. diameter and 1 m long silica tubes were purchased locally for drawing metal samples. One tube could be used for drawing 5 - 6 metal samples.

#### 5.2.6 Chemical reagents

HCl, HNO3, CdCl2, Ammonia solution used for chemical analysis of metal samples were of AR Grade, and Sodium thiosulphate, Potassium iodide were of GR grade. Iodine used was of LR grade.

#### 5.2.7 Argon gas

Argon gas cylinders were available locally.

#### 5.2.8 Standard steel sample

100 gr of standard carbon steel sample, supplied by British chemical standards No. 459/1 through their representative, Laboratory Chemical Co. Calcutta, in India, was used to calibrate the analysing equipment.

Complete description of the standard sample as reported by BSC is given in Table 5.2. Average values have been obtained by analysing the sample at many places using different methods. Manganese content varied from 0.95 to 0.99 pct. and sulphur content varied from 0.053 to 0.060 pct.

Table 5.2

Complete Analysis of the Standard Steel Sample as Reported by the BCS

									AND PROPERTY AND ADDRESS OF THE ADDR
Ele- ments			No.	of Ana	lysis				Ave-
Pct	1	2	2	4	5	6	7	8	rage
a	0.514	0.521	0.518	0.524	0.534	0,522	0.530	0.521	0.523
Si	0.580	0.590	0.560	0.580	0.570	0,580	0.580	-	0.580
Mn	0.960	0.950	0.960	0.950	0.980	0.960	0.990	0.980	0.970
P	0.052	0.051	0.053	0.056	0.057	0.054	0.052	0.054	0.054
S	0.056	0.053	0.059	0.058	0.060	0,058	0.056	0.054	0.057
Al	0.026	0.027	0.028	0.028	0.030	0.026		-	0.028
В	0.011	0.008	0.009	0.009	0.010	0.010	0.009		0.010
Q:o	0.105	0.105	0.108	0.111	0.108	0.111	0.116		0.109
Nb	0.017	0.012	0.013	0.014	0.015	0.013		***	0.014
Pb	0.002	0.002	0.002	0.002	0.002	garenty .	-		0.002
Λ	0.078	0.084	0.082	0.080	0.083	0.080	0.073	<del>-</del>	0.080
Zr	0.072	0.070	0.069	0.075	0.073			<del>grade</del>	0.072
Sb	0.006	0.006	0.006	0.007	0.006	0.006	0.006	_	0.006
									• 2*

#### 5.3 Experimental Procedure

#### 5.3.1 Calibration of thermocouple

The thermocouple (Pt-Pt + 10% Rh) was calibrated against melting point of pure copper, 99.9 pct.pure copper was taken in a graphite crucible and melted in the silicon carbide furnace under argon atmosphere. The arrest point was noted by recording temperature of the melt as function of time during cooling.

#### 5.3.2 Calibration of capillary flowmeter

The capillary flow meter was calibrated by noting the time of travel of the soap bubble in a glass tube of known volume, when argon gas was being passed, and by measuring at the same time the difference in heights in the two fluid columns of the flow meter.

#### 5.3.3 Making iron-carbon alloys.

The steel samples from Durgapur steel plant were machined to 35 mm dia. cylinders, which were then washed with acetone.

About 400 graphite prepared samples were put in graphite crucibles of 40 mm i.d., 10 cm long. The crucible with the steel sample was placed in the vacuum induction furnace. Steel samples melted in an hour or so and picked up carbon from the graphite crucible. The temperature of the furnace as recorded by the optical pyrometer was around 1550-1600 °C.

On cooling, the crucible was removed from the furnace and the alloy was recovered by breaking the crucible. It was then broken in to small pieces (10 mm size or less) by hammering. The sized Fe-C alloy was used subsequently for melting in the SiC furnace for investigations as described later.

#### 5.3.4 Making slags

Lime, alumina and silica materials were properly weighed and mixed to obtain the  ${\rm CaO/SiO}_2$  ratio 1, 1.5 and 2, with  ${\rm Al}_2{\rm O}_3$  percentage kept at 20 pct in the final slag.

The mixture was kept in a graphite crucible, 2.5 cm i.d. 7.5 cm long and placed inside the silicon carbide furnace under argon atmosphere at 1400°C. After 15-20 minutes, the constituents in the crucible became nearly molten. The molten slag was stirred by a graphite rod for half an hour. The cruicible was taken out of the furnace and allowed to cool. The solidified slag was obtained by breaking the cruicibles.

#### 5.3.5 Mn-S equilibrium in Fe-C alloys

The graphite crucible with 70-100 g of the Fe-C alloy and other additions such as Mn, FeS, MnS etc. was lowered into the reaction tube, which was being purged by argon before hand. The argon supply tube (6mm i.d. Al<sub>2</sub>0<sub>3</sub> tube) had to be taken out and reinserted into the reaction tube during the process of lowering the graphite crucible.

Continuous stirring of the melt was provided manually using a graphite rod. Sufficient time was allowed to reach equilibrium. The exact temperature of the melt was determined by placing the thermocouple just above the melt. Normally the temperature of melt differed from the temperature recorded by the other thermocouple by 30°C or so.

The alloy samples were sucked by dipping one end of the silica tube in the melt and using an aspirator at the other end. At a time 10-15 gm of samples could be collected. The sample tube was repeatedly purged with argon to minimise oxidation of melt by the entrapped air during sampling.

#### 5.3.6 Slag metal equilibrium

The graphite crucible was loaded with desired quantity of iron-carbon alloy and the premelted slag. The temperature was kept constant at 1400°C in such experiments to ensure that slag remained in the molten form. The alloy sample was sucked in similar fashion as described previously.

In some cases the crucible was taken out of the furnace after the experiment and samples were taken from the solidified mass after breaking the crucible.

# 5.3.7 Chemical analysis of samples

The samples to be analysed were first powdered to 100 mesh size in a pestle, mortar.

#### 5.3.7.1 Manganese

For analysis of Manganese by Atomic Absorption spectrophotometer the following steps were followed.

# 1) Preparation of solutions

0.1 gm of Mn (99.9 Pct pure), standard steel sample and unknown samples were weighed separately and dissolved in 10 ml of 1:3 HNO<sub>3</sub> and diluted to 100 ml after filtering. The stock solution prepared above were further diluted to the desired concentration. Solution of pure Mn were diluted to 0.2 ppm.

## 2) Procedure

The following instrumental condition for Mn analysis were fixed.

Hollow cathode lamp = Mn

Wave length = 279.5 nm

Flame = Air - acetylene

Absorbance of the solutions of known concentration is measured to obtain the linear plot between absorbance and concentration. From this plot and the measured absorbance of the unknown solution, its manganese concentration can be determined.

#### 5.3.7.2 Silicon

Some samples were analysed for silicon by gravimetric method in the Analytical Laboratory of Metallurgical Engineering Department, IIT Kanpur.

#### 5.3.7.3 Sulphur

Sulphur was analysed by evolution technique in the laboratory.

The following procedure was followed:

- (1) The set up was tested of any leakage of the gas. This was done by pressurising the vessel with argon and then noting any pressure drop in the line when the supply of argon was cut off.
- (2) Around 5 gm of the powdered sample was transferred carefully to the conical flask. The glass joint was placed on the proper position. The exit end of the glass tube was dipping in an ammonical cadmium chloride solution of 1 gm CdCl<sub>2</sub> in 60 ml of distilled water.

  80 ml of 1:1 HCl was poured into the flask from the separating funnel. The hot plate was switched on.

  The heat was regulated by operating the regulator of the hot plate. Heat supply was increased towards the end of the analysis in order to flush out the sulphur gas completely from the system by the vapours of H<sub>2</sub>O.

The sulphur in the alloy reacted with HCl to form  $H_2S$  gas, which was absorbed by the ammonical  $GdCL_2$  solution to form precipitate of cadmium sulphide. When it was confirmed that the alloy sample dissolved completely and  $H_2S$  gas was flushed out of the flask the absorbing solution was cooled in an ice bath. The ice cold solution was acidified with 20 ml of concentrated HCl. To the acidified solution 10 cc of N/10 iodine solution was added and titrated against N/10  $Na_2S_2O_3$  solution. 10 cc of N/10 iodine solution was also titrated against N/10  $Na_2S_2O_3$  solution in each case. The difference in two values of N/10  $Na_2S_2O_3$  amounts is the equivalent amount of  $Na_2S_2O_3$  solution consumed for sulphur in the sample.

The pct sulphur in the sample was calculated as follows:

$$\%S = \frac{0.16 \text{ Y}}{\text{X}}$$

where 'Y' is the amount of N/10  $Na_2S_2O_3$  solution consumed for sulphur and X is the weight of sample in grams.

# 5.3.8 Characterisation of phases by x-ray diffractometer

Prepared sample of less than 0.15 mm size was used for characterisation of phases of the materials.

The instrumental conditions were as follows: 20 angle was varied from 15° to 100°. The scanning speed was kept at 3° per min.

Cr-radition with  $\lambda = 2.291$  Å was used. From the peaks at different 20 values, the d values are calculated by the following relation;

$$\lambda = 2 \text{ d sin } \theta$$
.

#### CHAPTER VI

#### RESULTS

#### 6.1 <u>Fe-C-Mn-S System</u>

The desired quantity of iron carbon master alloy, manganese and sulphur in the form of FeS were added in the graphite crucible so as to give 100 gm of the liquid alloy on melting. In some cases, when sulphur in the form of MnS or FeS + MnS was also added to the graphite crucible, about 75 gm of Fe-C alloy was added to it. Table 6.1 summarises the quantity of materials added before or during melting.

# 5.1.1 Cooling and heating of melt at almost constant Mn level

The results of chemical analysis i.e. S and Mn content of the samples drawn at different temperatures from the melt in each case are also given in Table 6.1. The samples at about 1300°C were taken both during cooling and heating cycles in all cases to check whether equilibrium is reached or not.

In each case the melt was allowed to remain at 1300°C for one hour. In one experiment i.e. H<sub>6</sub> the melt was allowed to remain at 1300°C for another hour before drawing any additional sample. There was good agreement between the two values at the same temperature and this also signified the attainment of equilibrium in one hour at a particular temperature.

Table 6.1

Experimental [Mn] pct and [S] pct in Iron-Carbon Alloys at Temperatures  $1200^{0}$ - $1400^{0}$ C

Heat	Time Hours	Inpu	Input Materials	rials		Temp.	Holding time	Sample No.	Mn Nn 100 t.	S TO THE PERSON OF THE PERSON
		Fe-C Alloy	Mn	TeS	MnS	o .	moq		) )	거 ) •
	2		4	5	9	7		6	10	MACCO REALISTANT - ANTIBODANA REGISTRAÇÃO PROPRIADA PROPRIADA TANDOS POR TOTAL POR CONTRACTOR PO
Н	0.0	97.8	0.0	2.2	0.0	1360 1260		H-S- H-S2	N. A. N. A.	0.770 0.750
Н2	0- w4 0ror	98,111	-111	0.825	0.111	1400 1300 1200 1300	₹~~ <del>{</del> ~~ <del>{</del> ~~ <del>{</del> ~~ ~~	HH 2H 2	0.84 0.84 0.84	0.148 0.075 0.050 0.070
$_{\rm H_3}$	0.0 0.0 0.0 0.0	92.8	-111 8	1.04	0.0	1360 1290 1200 1290		H 2 1 2 1 2 1 2 1 2 1 2 1 2 2 2 2 2 2 2	0.91 0.91 0.90 0.90	0.116 0.087 0.047 0.084
$_{4}^{\mathrm{H}}$	0.5.04 0.00.00	96.5	Z • 1 1 1	1,26	0.111	1350 1290 1210 1290		НННН 444 4144 8888 -888	7.1.1 5000 5000	0.056 0.042 0.028 0.040

Table 6.1 contd...

	4.40			,		
- Company of the Comp	0.057 0.032 0.020 0.030	0.031 0.026 0.025 0.025 0.016	0.070 0.045 0.031 0.042	0.140 0.084 0.044 0.082	1.08	
10	0000 0000 0000	00000	00000	-000		
6	H5-81 H5-82 H5-83 H5-83	H H G L S L S L S L S L S L S L S L S L S L	Н Н Н Н Н Н В Н Н В В Н В В В В В В В В	Н Н Н Н Н Н Н Н Н Н Н Н Н Н Н Н Н Н Н	H9-S1	
8	4- 4- 4-	4- 5- 5- F- F-	~~~~	4004	-	
7	1360 1290 1200 1290	1360 1290 1290 1290	1400 1300 1300 1300	1400 1300 1300	1300	
9	0.0	0.1111	0.1111	**0*/	0.0	
5	.111	• 1 1 1 1	1.15	00.00	2.00	
4	K. 111	4.42	25.11.1	1.05	2,00	
3	94•0	400.4	95.0	70.0	50.0	
2	0.0 0.7 0.0 7.0	0-44v 0vvcv	0 + w 4 rv 0 rv 0 rv rv	0.0 4.5 7.0 9.5	0.0	
-	$^{ m H_5}$	岩	*#	$^{ m H_8}$	Н,	

\* 2.2 gms FeSi added at zero hour \*\* Iocal grade

Additional experiments were carried out to study the approach to the equilibrium during cooling of the melt without any stirring as to simulate the conditions in the ladle during the transit from the blast furnace section to the steel making section. Samples were drawn during cooling as soon as the temperatures of about 1300°C and 1200°C were recorded. The values are summarised in Table 6.2.

In one experiment i.e. H<sub>9</sub> enough Mn and FeS was added with the Fe-C alloy. After melting and holding for one hour the graphite crucible containing the melt was taken out and allowed to cool in air. It was then broken to collect the metal and slag samples for chemical analysis and X-ray diffraction study.

## 6.1.2 Stepwise additions of manganese at constant temperature

In some experiments Mn content of the melt was increased by stepwise addition of Mn at a constant temperature. The results of these experiments are summarised in Table 6.3.

# 6.2 Slag Metal Equilibrium Studies

The desulphurisation capacity of the oxide slag S' was studied at different manganese levels in the metal at 1400°C. Mn was added to the melt in steps and samples were drawn through silica tube after allowing the melt to reach equilibrium in 4 hours with constant stirring of the bath. The details of experiments and the results of chemical analysis are given in Table 6.4.

Table 6.2 Results of Experiments of the Cooling the Melt from  $1400^{\circ}\text{C}$ 

Heat No.	Sample No.	Input	materi	als,gms	Time	Temp.	Metal	Analysis
110.	140	Fe-C Alloy	Mn	FeS	Mins.	°G	Mn	pct S
	H <sub>10</sub> -S <sub>1</sub>	80	1.9	0,9	0	1390	1.7	0.083
H <sub>10</sub>	H <sub>10</sub> -S <sub>2</sub>	pura.	-	timb.	10	1275	1.7	0,048
	H <sub>10</sub> -S <sub>3</sub>		,		15	1220	1.7	0.044
	H <sub>11</sub> -S <sub>1</sub>	80	1.9	0.9	0	1388	1,6	0,065
H <sub>1</sub> 1	H <sub>11</sub> -S <sub>2</sub>	pope	***	-	44	1300	1.6	0.038
	H <sub>11</sub> -S <sub>3</sub>	-		-	84	1220	1,6	0.031

Table 6.3

Experimental Results of Stepwise Addition of Mn at  $1325^{\rm o}$  and  $1400^{\rm o}$ 

	10d	0.084 0.047 0.036	0.347 0.110 0.069 0.057	0.680 0.230 0.095 0.064
To the	a od	1.27 2.35 3.26	2.771 2.780 3.140	0.20 0.600 1.320 160
Sample	отпО * о ь	17.TO	0000	10 10 20 20
Sumple	• 01-	H 2 81 H 2 82 H 2 82	н н н н н г г г г г г г г г г г г г г г	HEHH 141 141 150 150 150 150 150 150 150 150 150 15
Holding	Hour	N	4 4 2	4400
remp.	٥ 0	1325 1325 1325	1400 1400 1400 1400	1400 1400 1400
errikan dan dan dan salam dan s	MnS	0,0	0.0	* * * * * * * * * * * * * * * * * * *
13 1 gms	168	6.875*	**000.7	* * * * * * * * * * * * * * * * * * * *
Input Marerials	Re-G Wn	0.1 0.0	0.00	0.00
Heav Time Input Marerials	re-d Alloy	89.4	70.0	70.0
mime	Jnou	0 0 <b>m</b>	2 <del>4</del> 8 0	0480
Hear		H 12	H 2	H 4

Imported grade Iocal grade \*

<sup>\*\*</sup> 

dentd....

Table 6,4

Experimental Results of Slag-Metal Equilibrium at 1400°C

		}		,	Ber-ridger Ber-		
	Sipot	<del>-</del>	1	1 1	esperantes est de la constante est de la const		
	To the control of the	,			general control of the control of th		
	Pod Foot	10	0.256	0.152	0.108	0.070	
· · · · · · · · · · · · · · · · · · ·	andraides (Park V - 2 -		-		**************************************		
<b>1</b>	Mn pet	6		1 1	1,20	1.15	
120°3 20°3 20°3 20°3	Property of the state of the st					er de seguine de la company de	
	Sample No.		H <sub>15</sub> -S <sub>1</sub>	H <sub>16</sub> -S <sub>1</sub>	H <sub>17</sub> -S <sub>1</sub>	H <sub>18</sub> -S <sub>1</sub>	
Sic 2 40 22 26.6		Carrier and Company of the Company o		The state of the s		To the second control of the second control	
	Hold time bour	7	ત ત	2 2	2 2	2	
4	lag anti	9	4-4	8 1	<b>R</b> -4	8	
Gao 440 57.44	erials Mr	5	00.0	O O O	1 and a distribution of the 20	1.20	
	Input Materials gms d res Mn	4	1,00	r univalina adronis st 1 ,00	0,90	0.90	Marian Carallana Carana
Slags		3	75.0	80.0	78.8 7	78.0	e filmeline Tabilité de cerrer
	Time Hour	2	0 8	0 7	2	0 2	A Security of the
	Heat 10.	** ***********************************	H15	H16	1117	H <sub>18</sub>	And the same of the same of the same

Table 6.4 contd....

¥	j.	1	į	σ <sup>†</sup>	3	<del></del>
11		er green jan de gewenne gewenn		110	1 1 1	0.31
10	0.073	0.054	0.116	0.096 0.070 0.040	0.076 0.048 0.035	0.046 0.028 0.016
6	2.07	2.20 2.12	1.25	1,00	1.09	1.27
	H19-31	H20-S1 H20-S2	H21-31 H21-32 H21-32	H22-51 H2 <b>2-</b> 52 H22-53	H23-31 H23-32 H23-52	H24-51 H24-52 H24-53
	22	22	444	444	44	444
6 Sharet a selection and selec	4-4	14-8	4-15	Andreas and the second of the	3rd 7.08	B-15
To the same of the	2.0	2.10	0.00	09.0	0.00	0.00
4	0.87	06.0	0.83	0.834	0.830	0,820
and support Manay and Later Contracts of Con	78.0	73.0	75.0	75.0	0 * 9 1	75.0
2	0	0 2	0 4 8	0 4 8	0 4 8	0 0 4 8
The section of the se	H <sub>19</sub>	H20	H21	H22	**H23	†2H

Contd. ...

Table 6.4 contd...

_	α	κ	4	ĸ	•	<b>t</b>	80	o,	10	
10 73 11	040	125 4	0.83	0.00	0-7.9	444	H25-51 H25-52 H25-53	1.23	000	0.060
H26	O4 Ω	H <sub>26</sub> 0 75.0 0.83	0.83	0.00 0.98 0.60	0 15	444	H26-51 H26-52 H26-52	1.31	0.0	0.009 0.009 0.006
i Li	<b>⊕</b> → ∞	76.0	0.9	0.00	7.7-0	444	H27-S1 H27-S2 H27-S3	1.29	0,010 0,008 0,007	10 08 07

\* 1.61 pot Mn in final slag \*\* 0.76 pot Mn in final slag \*\*\* 0.54 pot Mn in final slag

Table 6.5

•	Experime	Experimental Results	ults to	Esta	blish Sla	Establish Slag-Metal Equilibration	ui li br <b>ati</b>	on Time at	Time at 1400°C	
			Slag		GaO	Sio		A1,002		
			<b>ৰ</b> চ		53.4	40 26.6		2002		
Heat No.	Time Hour	Inpu	Input Materials in gms	ials i	In gms	Holding time	Sample	Mn	CC 'Annual transmission of the Control of the Contr	A. American
		Fe-C Alloy	FeS	Mn	Slag- Quantity	hour	•	3 2 2 7	poct	
þ	0 -	75.0	0.83	1.21	A-15	4	H <sub>28</sub> -S <sub>1</sub>	1.29	0,068	
п28	4 00	1 1	1 1	1 1	1 1	4 4	H <sub>28</sub> S <sub>2</sub> H <sub>28</sub> S <sub>3</sub>	1.20	0.067	
	0	75.0	0.82	0.00	(3-7.5		n n	ANTACA GALLA C'ANTRONOMINA COMPONINA DE LA COM	ARTHUR RECEPTANT AND THE ARTHUR ARTHU	- 14
H <sub>29</sub>	4	ı	4			+ 4	H29-52	ŀ [	0.049	
·	σ	¥ .	i	1	ī	4	H <sub>29</sub> -S <sub>3</sub>	i	0.050	

Table 6.6

Experimental Results of Slag Metal Equilibrium at 1400°C

H <sub>30</sub> H <sub>30</sub> -S <sub>1</sub> 30 0.33 H <sub>31</sub> H <sub>31</sub> -S <sub>1</sub> 30 0.33 H <sub>32</sub> H <sub>32</sub> -S <sub>1</sub> 30 0.33 H <sub>34</sub> H <sub>34</sub> -S <sub>1</sub> 30 0.33 H <sub>36</sub> H <sub>35</sub> -S <sub>1</sub> 30 0.33 H <sub>36</sub> H <sub>35</sub> -S <sub>1</sub> 30 0.33 H <sub>36</sub> H <sub>35</sub> -S <sub>1</sub> 30 0.33 H <sub>36</sub> H <sub>36</sub> -S <sub>1</sub> 30 0.33 H <sub>39</sub> H <sub>39</sub> -S <sub>1</sub> 30 0.33 H <sub>40</sub> H <sub>40</sub> -S <sub>1</sub> 30 0.33	it materials	in gms		Compound	ਲ	Holdin	Wetal	Analysis
H <sub>30</sub> -S <sub>1</sub> 30 0.33 H <sub>31</sub> -S <sub>1</sub> 30 0.33 H <sub>35</sub> -S <sub>1</sub> 30 0.33 H <sub>35</sub> -S <sub>1</sub> 30 0.33 H <sub>36</sub> -S <sub>1</sub> 30 0.33 H <sub>36</sub> -S <sub>1</sub> 30 0.33 H <sub>36</sub> -S <sub>1</sub> 30 0.33 H <sub>39</sub> -S <sub>1</sub> 30 0.33 H <sub>40</sub> -S <sub>1</sub> 30 0.33		Slag	Gan %	Sio	A12.03	Hour	Mn	Hour Mn sot S
H <sub>31</sub> -S <sub>1</sub> 30 0.33 H <sub>32</sub> -S <sub>1</sub> 30 0.33 H <sub>34</sub> -S <sub>1</sub> 30 0.33 H <sub>35</sub> -S <sub>1</sub> 30 0.33 H <sub>36</sub> -S <sub>1</sub> 30 0.33 H <sub>37</sub> -S <sub>1</sub> 30 0.33 H <sub>39</sub> -S <sub>1</sub> 30 0.33 H <sub>40</sub> -S <sub>1</sub> 30 0.33		9	40.0	40.0	50	9		960*0
H <sub>32</sub> -S <sub>1</sub> 30 0.33 H <sub>34</sub> -S <sub>1</sub> 30 0.33 H <sub>35</sub> -S <sub>1</sub> 30 0.33 H <sub>36</sub> -S <sub>1</sub> 30 0.33 H <sub>36</sub> -S <sub>1</sub> 30 0.33 H <sub>39</sub> -S <sub>1</sub> 30 0.33 H <sub>40</sub> -S <sub>1</sub> 30 0.33		80	40.0	40.0	20	9	0.45	0.135
H35-S1 30 0.33 H34-S1 30 0.33 H36-S1 30 0.33 H36-S1 30 0.33 H38-S1 30 0.33 H39-S1 30 0.33 H <sub>0</sub> -S1 30 0.33		9	40.0	40.0	20	9	0,48	960.0
H <sub>3</sub> 4-S <sub>1</sub> 30 0.33 H <sub>3</sub> 5-S <sub>1</sub> 30 0.33 H <sub>3</sub> 6-S <sub>1</sub> 30 0.33 H <sub>3</sub> 7-S <sub>1</sub> 30 0.33 H <sub>3</sub> 9-S <sub>1</sub> 30 0.33 H <sub>4</sub> 0-S <sub>1</sub> 30 0.33	0.33 0.434	9	40.0	40.0	50	9	0.99	0.071
H <sub>35</sub> -S <sub>1</sub> 30 0.33 H <sub>36</sub> -S <sub>1</sub> 30 0.33 H <sub>37</sub> -S <sub>1</sub> 30 0.33 H <sub>39</sub> -S <sub>1</sub> 30 0.33 H <sub>40</sub> -S <sub>1</sub> 30 0.33		9	48.0	32:0	82	9	**	0.043
H <sub>36</sub> -S <sub>1</sub> 30 0.33 H <sub>37</sub> -S <sub>1</sub> 30 0.33 H <sub>39</sub> -S <sub>1</sub> 30 0.33 H <sub>40</sub> -S <sub>1</sub> 30 0.33		9	48.0	32.0	20	9	0.47	0.041
H <sub>37</sub> -s <sub>1</sub> 30 0.33 H <sub>38</sub> -s <sub>1</sub> 30 0.33 H <sub>40</sub> -s <sub>1</sub> 30 0.33		9	48.0	32.0	50	9	0.92	0.030
H <sub>3</sub> g-S <sub>1</sub> 30 0.33 H <sub>3</sub> g-S <sub>1</sub> 30 0.33 H <sub>40</sub> -S <sub>1</sub> 30 0.33		~	48.0	32.0	50	Q	0.55	0.075
H <sub>39</sub> -S <sub>1</sub> 30 0.33 H <sub>40</sub> -S <sub>1</sub> 30 0.33		Q	53.4	56.6	20	9	1	0.020
H <sub>40</sub> -S <sub>1</sub> 30 0.33		9	53.4	56.6	20	9	0.53	0.016
- 0#		9	53.4	56.6	20	9	1,08	0.011
H41-S1 30 0.33		20	53.4	26.6	8	9	0.47	0.017

Table 6.7
Results of Rechecked Sulphur Determinations on Metal Samples

Sample No.	Pet Sul	phur
and an interpretability of the control of the contr	1st Analysis	2nd Analysis
H <sub>3</sub> -S <sub>1</sub>	0.116	0.112
H <sub>4</sub> -S <sub>4</sub>	0.040	0.039
H <sub>5</sub> -S <sub>4</sub>	0.029	0.030
H <sub>17</sub> -S <sub>2</sub>	0.096	0.116
H <sub>21</sub> -S <sub>3</sub>	0.040	0.042
H <sub>24</sub> -S <sub>3</sub>	0.016	0.014

Table 6.8

Analysis of Standard Steel Sample in the Laboratory

Sl.No.	Percentage of S Reported, Average Value	Percentage of S As Analysed in the Laboratory
1	O 057	0.054
-	0.057	
2		0.051
3		0.048
4 .		0.049
5		0.052
6		0.053
7		0.048
8		0.051
LANGE THE STREET STREET AND THE STREET S	Average	0.051

# 6.4 X-Ray Characterisation of Materials

X-ray diffraction pattern to characterise various materials such as steel samples from Durgapur, Manganes (imported) iron sulphide (imported), iron sulphide (local), manganese sulphide (local), prepared iron-carbon master alloy in the laboratory by induction melting, prepared manganese sulphide in laboratory by the reaction between stochiometric amounts of molten iron sulphide and manganese in graphite crucible at around 1365°C, collected slag samples of some heats are given in Figures 6.1 to 6.4.

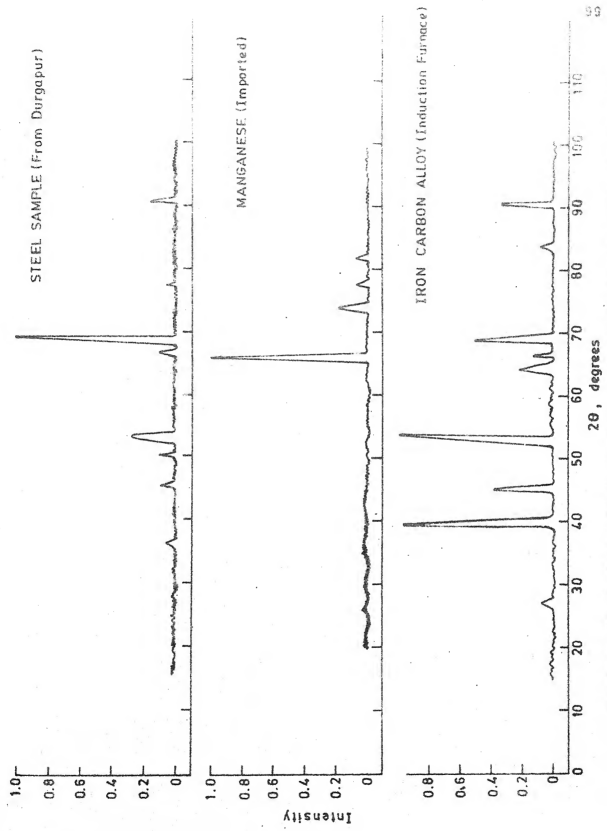
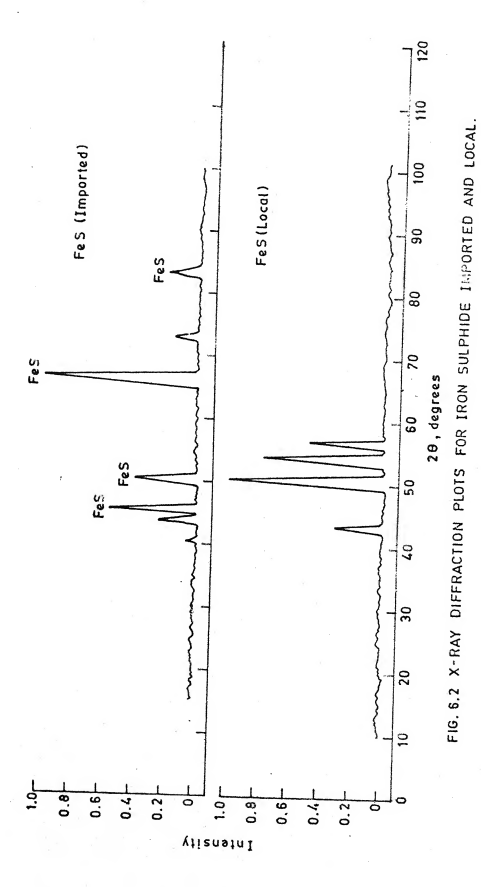
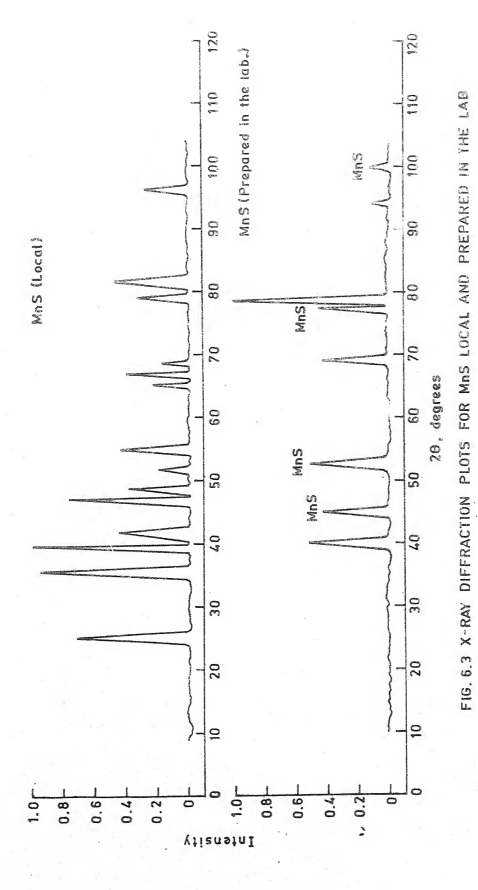
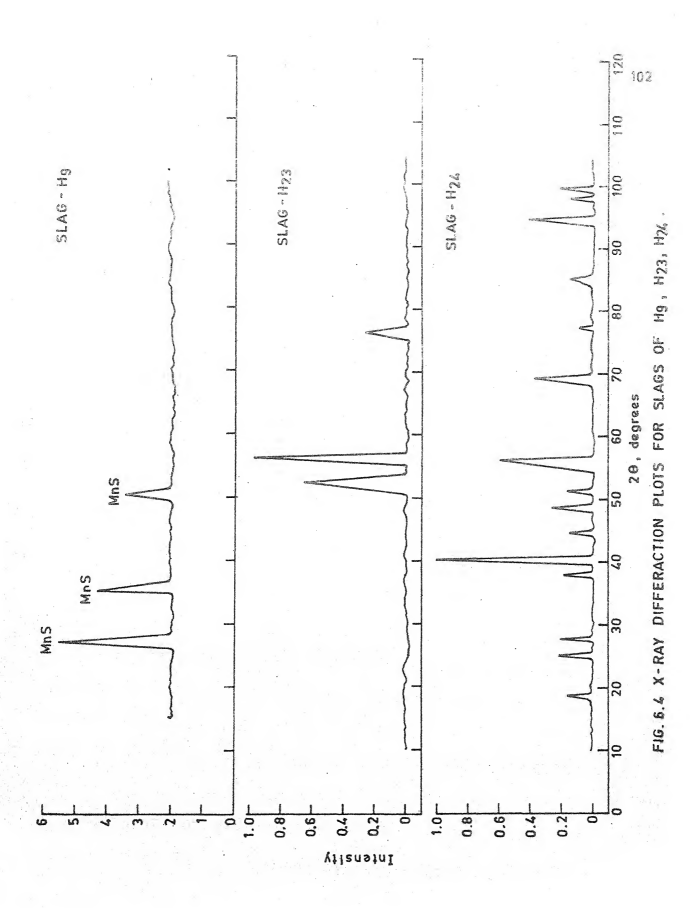


FIG. 6.1 X-RAY DIFFRACTION PLOTS FOR IRON, MANGANESE AND IRON-CARBON ALLOY







#### CHAPTER VII

#### DISCUSSION

## 7.1 Manganese Sulphur Equilibrium

# 7.1.1 Reliability of data

Determination of the manganese or sulphur contents in the samples of the iron alloys as reported in Chapter 6 can be believed to be quite accurate because of the good agreement obtained between the reported analysis and the actual analysis of the standard steel samples, and also because of the agreement reached between the two values of the same sample in a few cases. For these melts quantity of the slag formed may be around one percent or so and this did not pose any problem in sucking the metal sample from the bottom of the crucible. From the theoretical analysis given in Chapter 4, it is possible to calculate the equilibrium sulphur level in the melt at a particular temperature and manganese content of the metal. The experimental values of the sulphur analyses are compared with the calculated values in Figure 7.1.

Agreement is quite good especially for those samples which were taken at  $1200^{\circ}$ C. Similar analysis was done to compare the experimental values of Morris<sup>14</sup> with the calculated values as shown in Figure 7.2. Morris's <sup>14</sup> results agreed with the theoretical calculations both at  $1200^{\circ}$ C and  $1400^{\circ}$ C. At higher temperatures of  $1350^{\circ}$ C and  $1400^{\circ}$ C, the experimental results of the present study are lower than the calculated values. This discrepancy may be attributed due to any of the

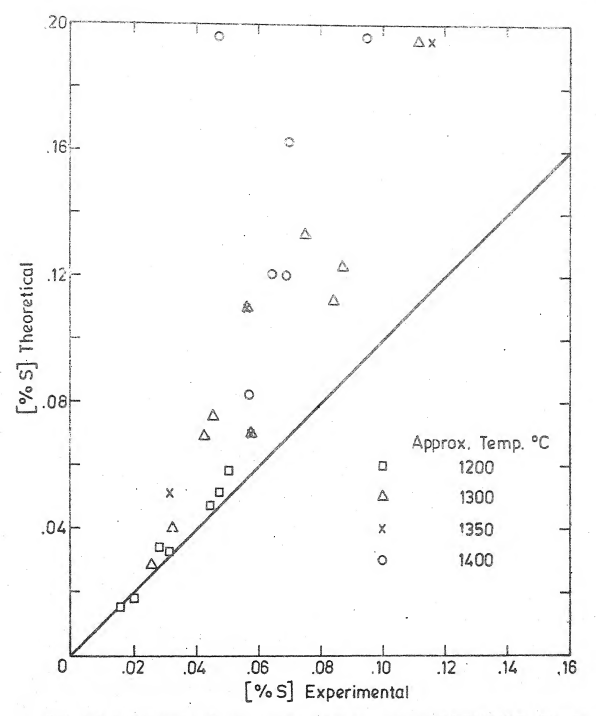


FIG. 7.1 COMPARISON OF EXPERIMENTALLY DETERMINED SULPHUR VALUES WITH THEORETICALLY CALCULATED SULPHUR VALUES FROM THERMODYNAM DATA.

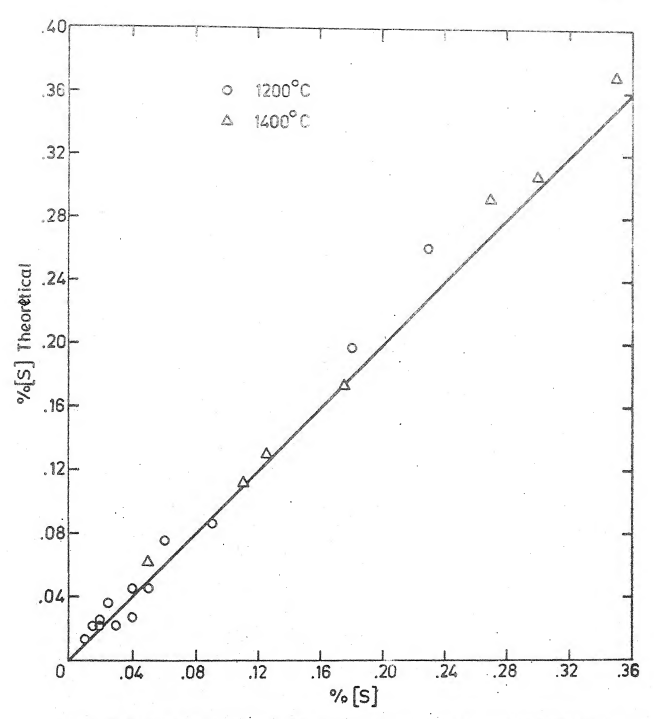


FIG. 7.2 COMPARISON OF MORRIS'S DATA WITH THEORETICALLY CALCULATED SULPHUR VALUE

### following;

- (1) Experimental errors in estimating temperature, manganese and sulphur levels of the melt.
- (2) Formation of MnO, FeO or SiO<sub>2</sub> in the slag so as to lower the activity of MnS in the slag at higher temperatures.
- (3) Formation of FeS in the slag especially at higher sulphur levels of the melt as to lower the activity of MnS in the slag.

Experimental errors in estimating temperature, manganese or sulphur can not explain the discrepany because enough care was taken to control them. Thermocouples were properly and periodically calibrated and the tip of the thermocouple lied very close to the melt. The argon tube was positioned at 20 cm above the melt and it cannot account for any temperature gradient within the melt.

Any FeO or MnO formed due to oxidation of metal by oxygen in the argon gas will react immediately with graphite crucible as per the following reaction;

$$(FeO) + C(gr) = Fe + CO$$
 (7.1)

$$(Mn0) + C(gr) = [Mn] + C^{\circ}$$
 (2.31)

$$K_{2.31} = \frac{f_{Mn} [Wt\% Mn]}{a_{Mn0} a_{C}} p_{CO}$$
 (7.2)

$$\log K_{2.31} = -\frac{15090}{T} + 10.90 \tag{7.3}$$

For carbon saturated iron  $a_{C} = 1$  and  $f_{Mn} = 0.8$  and taking  $p_{CO} = 1 \text{ atm.}$ , from equation (7.2) and (7.3),  $a_{MnO}$  value at 1300°C is 0.039 for one percent manganese in the melt. Since reaction is occuring at the slag metal interface  $p_{CO}$ can be taken as one atm. Otherwise its value will be lowered as to decrease  $a_{\overline{\text{MnO}}}$  further. Thus thermodynamic calculations show that  $a_{\text{MnO}}$  in the slag at 1300°C cannot be greater than 0.05. Similarly FeO in the slag will be negligible. oxidation of metal can be ruled out. As steel samples were obtained at the end of refining in the open hearth steel making, presence and oxidation of any silicon in the melt to give SiO2 in the slag is also very unlikely. There can not be any chance of entrapped slag coming with the steel samples as they were first machined and remelted in the induction furnace to make the iron-carbon alloys.

Both manganese and sulphur are dissolved in the melt at higher temperature and as the temperature is lowered the equilibrium product of [Mn][S] is lowered. Manganese and sulphur must react to form MnS which then floats to the surface. Mucleation of MnS inside the melt is essential for reaching equilibrium. Similarly iron and sulphur may react to form FeS, if the activity of FeS be lowered by its solution in MnS. Exact behavior of FeS - MnS system is not known, but if it is assumed to be ideal, a<sub>MnS</sub> is likely to be close to unity. Formation of FeS can thus not explain the observed lowered values of sulphur in the melts compared to the theoritical values, especially in the melts having sulphur content less than 0.06 percent.

It can be concluded from the above that the consistent lower values of sulphur obtained at 1300°C and 1400°C compared with the theoretical values may be attributed to the higher activity coefficient of sulphur in the melt or to the higher equilibrium constant for the reaction (4.5), than the values reported in the literature. The equilibrium constant and the standard free energy of the reaction (4.5) are recalculated using the experimental data of the present investigation and the activity coefficient values of sulphur and manganese as calculated in Chapter 4. The calculated values of  $\triangle G^{0}$  are compared with the reported values from equation (4.6) in Figure 7.3. The discrepancy is greater for samples taken at temperatures of 1300°C and 1400°C compared to these samples taken at 1200°C. At higher temperature, sulphur content is more which can lead to the formation of FeS in the slag. Probability of FeS and MnS to form solid or liquid solution is also more at higher temperatures. By taking amns as mole fraction of MnS in the slag as explained in Chapter 4 the reported values of AGO agree better with the recalculated values from the experimental data of the present study. In most of the cases at 1300°C and 1400°C, discrepancy in the two values of AGO is less than 1.8 kcal/mole which is within the accuracy of the reported data.

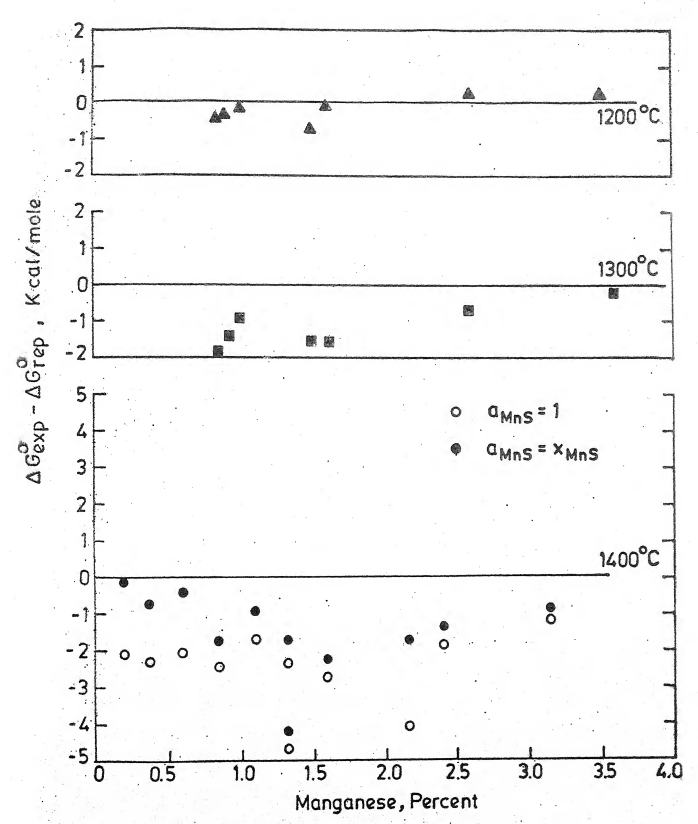


FIG. 7.3 PLOT SHOWING DISCREPANCIES IN THE EXPERIMENTALLY DETERMINED VALUES OF STANDARD FREE ENERGY OF FORMATION OF MnS FROM [Mn] AND [S] IN MELT.

# 7.1.2 Cooling and heating of the melt at the same manganese level

The sulphur analysis of the samples collected during cooling to 1200°C and then heating back to 1300°C revealed that there was some sulphur transfer from the slag to the metal during the reheating period. The sulphur level in such samples was always found to be lower than that of the samples taken during cooling of the melt. This discrepancy may be due to the fact that the sulphur transfer from the slag to the metal is sluggish as the quantity of MnS formed is very small and it may not be uniformly mixed with the melt during the mannual stirring by the graphite rod. The dissolution kinetics of MnS is not well understood but it may take more than one hour at 1300°C to reach equilibrium with the melt. The fact that MnS in slag decomposes to approach equilibrium between the Mn and S levels in the melt

# i.e. MnS = Mn + S

is confirmed by the results of the heat Hg, in Table 6.1.

Theoritically it is known that manganese and sulphur in the melt will react to form MnS in the slag as to reach equilibrium. However the quantity of slag formed was so small that it could not be collected and analysed easily. The quantity of slag formed in one case was deliberately increased by adding excessive amount of FeS and Mn to the charge materials in the crucible of heat Hg. In this case the slag could be collected after breaking the crucible at the end of the experiment. Comparison of X-ray diffraction

of this slag(Figure 6.4) with the X-ray diffraction file data confirmed the presence of MnS. Presence of FeS or Fe phases in slag was not noticed. Chemically it was analysed to contain more than 35 pct. Mn and 20 pct S. The set up was not geared to analyse such samples accurately so the results can be subject to some errors. These findings however confirmed the formation of MnS in the slag. This is also confirmed from the results reported in Table 6.3 that sulphur in melt decreases when manganese additions to melt at a constant temperature. Attempts were also made to form MnS in the laboratory by melting FeS and Mn materials in the graphite crucibles. X-ray diffraction pattern of the product Figure 6.2 marked the presence of MnS and some other phases. X-ray diffraction patterns of locally procurred FeS and MnS materials however did not give any clear picture regarding the phases present in them. Exact chemical analysis of these materials are also not known though they were analysed to contain about 20 to 30 pet sulphur.

# 7.1.3 Approach to the equilibrium during furnace cooling of the melt

Attempts were made to simulate the plant conditions where metal in the ladle cools without any external stirring. When the melt was allowed to cool in the furnace under the argon atmosphere by simply turning off the power of the furnace, it took 10 minutes for the melt to reach 1300°C and another 5 minutes to reach around 1200°C. The sulphur analysis of the samples collected from the cooling melt is compared with the sulphur analysis of the samples from

the melt where equilibrium was allowed to reach by holding the melt at that temperature for one hour or so, in Figure 7.4. The temperature measurements of the cooling melt may be slightly erroneous but the trend is towards higher S values in the samples from the cooling melt. Still a significant drop in the sulphur content during the cooling of melt in 10-20 minutes of time is observed and it may be quite significant from practical point of view. The sulphur levels tend to reach the equilibrium values if the melt is cooled slowly in the furnace by reducing the power to the furnace such that it takes 40 and 80 minutes for the melt to reach 1300°C and 1200°C respectively.

# 7.1.4 Model for calculating manganese and sulphur in the melt

A mathematical model based on manganese and sulphur balances can be worked out to determine the manganese and sulphur contents of the melt under certain equilibrium conditions. Let the quantities of FeS, MnS and Mn, in gms added to 100 gms of Fe-C alloy be expressed as  $X_1, Y_1$  and  $Z_1$  respectively. If  $X_2$  gm of FeS and  $Y_2$  gm of MnS decompose to give Fe, Mn and S to the melt, the final quantity of the molten alloy will be  $(100 + X_2 + Y_2 + Z_1)$ . Similarly slag quantity will be  $(X_1 + Y_1 - X_2 - Y_2)$  gms and percentage of manganese and sulphur in the molten alloy can be written as follows;

$$[Mn] = \frac{55}{87} \frac{Y_2 + Z_1}{100 + X_2 + Y_2 + Z_1} \times 100$$
 (7.4)

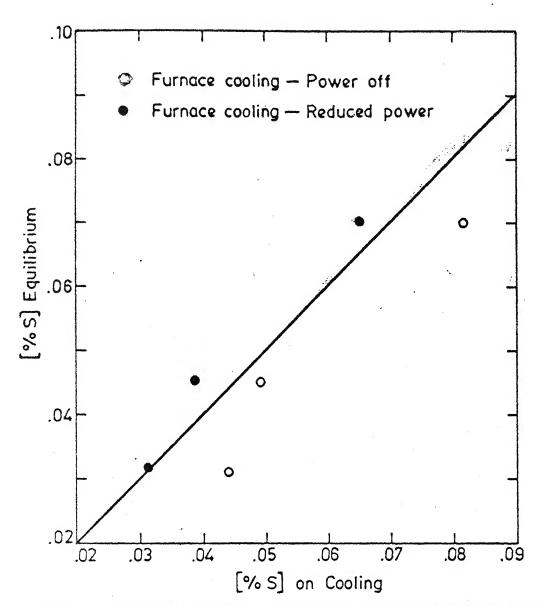


FIG. 7.4 PLOT SHOWING APPROACH OF EQUILIBRIUM DURING COOLING FROM 1400°C.

$$[S] = \frac{32 \frac{X_2}{88} + \frac{Y_2}{87}}{100 + X_2 + Y_2 + Z_1} \times 100$$
 (7.5)

Assuming ideal behavior of FeS and MnS in the slag, the equilibrium constants for reactions (4.5) and (4.15) can be written as follows:

$$K_{4.5} = \frac{X_{MnS}}{f_{Mn} f_{S}[Mn][S]}$$
 (7.6)

$$K_{4.15} = \frac{X_{\text{FeS}}}{f_{\text{S}}[s]}$$
 (7.7)

The mole fraction of MnS and FeS can be assumed to be same as their weight fraction and can be written as follows;

$$x_{\text{MnS}} = \frac{y_1 - y_2}{x_1 + y_1 - x_2 - y_2}$$
 (7.8)

$$x_{\text{FeS}} = \frac{x_1 - x_2}{x_1 + y_1 - x_2 - y_2}$$
 (7.9)

Values of  $K_{4.5}$  and  $K_{4.15}$  at any temperature can be known from the data given in Chapter 2 or 4. To compute the values of  $K_2$  and  $K_2$  equations (7.6) to (7.9) can be solved. Then the percentage of mangane and sulphur in the melt can easily be computed by using equations (7.4) and (7.5).

For example using experimental data of heat  $\rm H_9$  manganese level in melt was calculated to be around 0.1 pct and the sulphur level as 1.17 pct from the above model. The calculated value agreed with the experimentally determined sulphur content of 1.07 pct. Similarly for heats  $\rm H_1$  to  $\rm H_7$ 

calculations showed that manganese should react with added FeS to form MnS and the final Mn in the alloy would be 0.2 to 0.6 pct less than the value calculated from the manganese additions to the charge. In case of heats H13 and  $H_{14}$  local grade of MnS and FeS were used, the [Mn] and [S] levels calculated from the above model did not match well with the experimental values as shown in Table 7.1. only source of manganese and sulphur in the alloy melt was MnS in heat : H13 as no manganese or FeS was added in the charge mix of the crucible. Some MnS might decompose to give manganese and sulphur in the alloy melt and some MnS might react with Fe to form FeS and manganese. Thus the ratio of manganese and sulphur in the alloy melt should be greater than the stoichiometric ratio of 55:32 where as actual Mn:S ratio of 1.06 was obtained. This was possible if some sulphur was present in some other form e.g. MnS2, FeS, free sulphur etc. in the charge material.

# 7.2 Slag-Metal Equilibrium

# 7.2.1 Effect of slag basicity on sulphur levels of the melt

The effect of slag basicity for manganese free metal on its sulphur level is plotted in Figure 7.5 based on the experimental data of Chapter 6. As the basicity of the slag i.e. (CaO/SiO<sub>2</sub> ratio) increases, the desulphurisation of the metal increases. This agrees with the findings of many other investigators. Increasing the slag quantity has also helped in decreasing the sulphur level of the melt. Corresponding calculations for the sulphur partition ratio and comparison

Table 7.1

Comparison of Theoritical and Experimental [Mn] and [S] Levels at 1400°C

Heat	Sample No.	pct. Mn	ON THESE WINDS AND	pet. S	AND AND SOME OF THE PARTY AND ADDRESS OF THE PARTY ADDRESS OF THE PARTY AND ADDRESS OF THE PARTY ADDRESS O
COMMUNICATION THE COMMUNICATIO	-10 4	Theoritical	Expt.	Theoritical model a <sub>Mns</sub> = X <sub>MnS</sub>	Expt.
	H <sub>13</sub> -S <sub>1</sub>	0.955	0.371	0.188	0.347
TT	H <sub>13</sub> -S <sub>2</sub>	1.910	1.320	0.099	0.110
<sup>H</sup> 13	H <sub>13</sub> -S <sub>3</sub>	2.840	2.160	0.065	0.069
	H <sub>13</sub> -S <sub>4</sub>	2.760	3.140	0.051	0.057
Marie Casa Nazari erinasus — eta esta 31 cipilosofia 2	H <sub>14</sub> -S <sub>1</sub>	0.162	0.200	0.734	0.680
H <sub>14</sub>	H <sub>14</sub> -S <sub>2</sub>	0.350	0.600.	0.450	0.230
14	H <sub>14</sub> -S <sub>3</sub>	0.660	1.320	0.276	0.095
	H <sub>14</sub> -S <sub>4</sub>	1.150	2.160	0.160	0.064

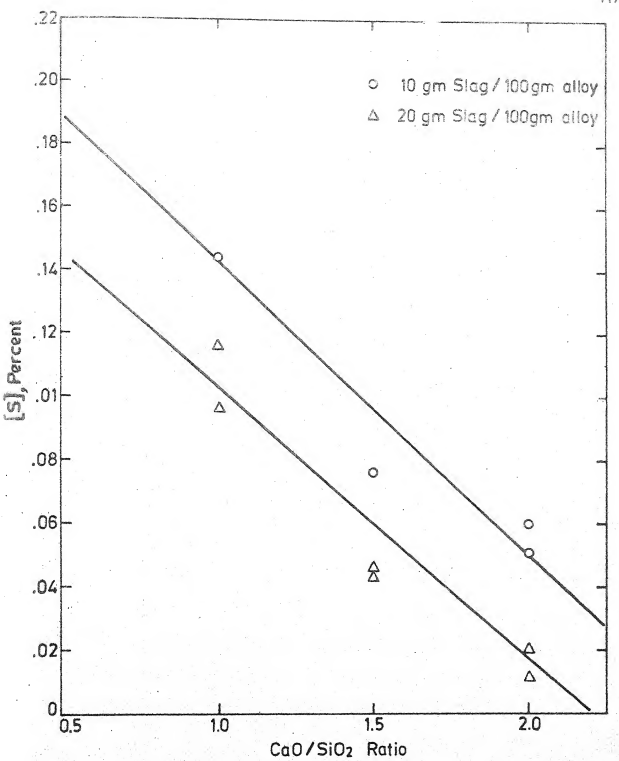


FIG. 7.5 EFFECT OF SLAG BASICITY (CaO/SiO2 ratio) ON THE DESUL-FURISATION OF CARBON SATURATED IRON AT 1400°C.

with the theoritical value are described later.

# 7.2.2 Effect of manganese on sulphur levels of the melt

At 1400°C as the manganese level in the metal increases, the sulphur level decreases in almost all cases as shown in Figures 7.6 to 7.9 for different compositions and volumes of the slag.

It has been discussed in Chapter 4 that the sulphur level of metal should decrease with decreasing a<sub>MnS</sub>. MnS formed in presence of Mn in the melt will lower its activity by becoming a part of the slag. Transfer of manganese from metal to slag phase was confirmed by the chemical analysis of the slag in a few cases. Actual transfer of manganese from metal to slag can result due to the formation of MnS or MnO as per equations (4.5) and (2.34). X-ray diffraction of the slag (Figure 6.4) did not confirm the presence of MnS or MnO phases probably due to small quantity of MnS or MnO in the slag or because they may be present in some other form. The role of manganese in lowering the sulphur level can be attributed to any of the following;

- (1) Manganese reacts with sulphur to form MnS and activity of MnS is lowered by its solution with the oxide slag.
- (2) Manganese enhances the rate of sulphur removal by CaO in the slag as per the following reactions;

$$(cao) + \underline{s} = cas(s) + \underline{o}$$
 (7.10)

$$0 + Mn = (Mn0)$$
 (7.11)

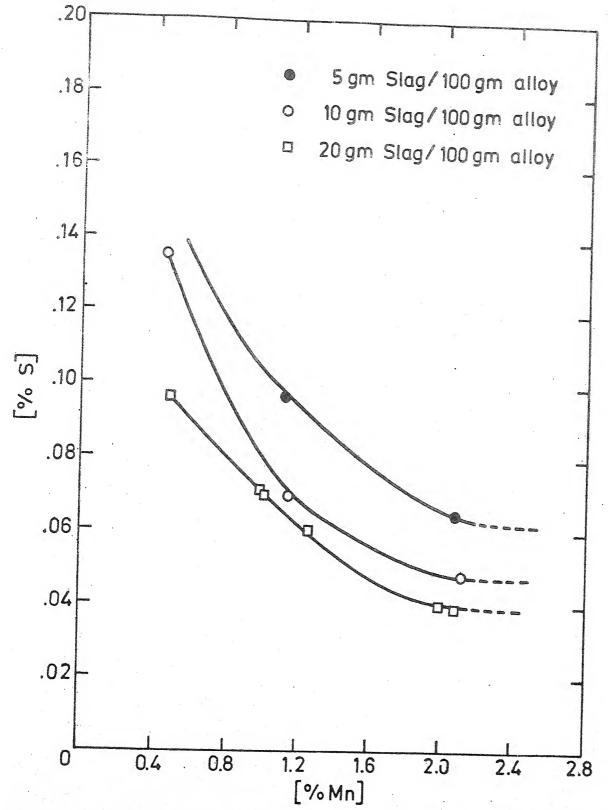


FIG. 7.6 EFFECT OF MANGANESE IN CARBON SATURATED IRON ON THE DESULPHURISATION OF THE MELT AT 1400°C AND SLAG BASICITY (CaO/SiO2 Ratio) OF ONE.

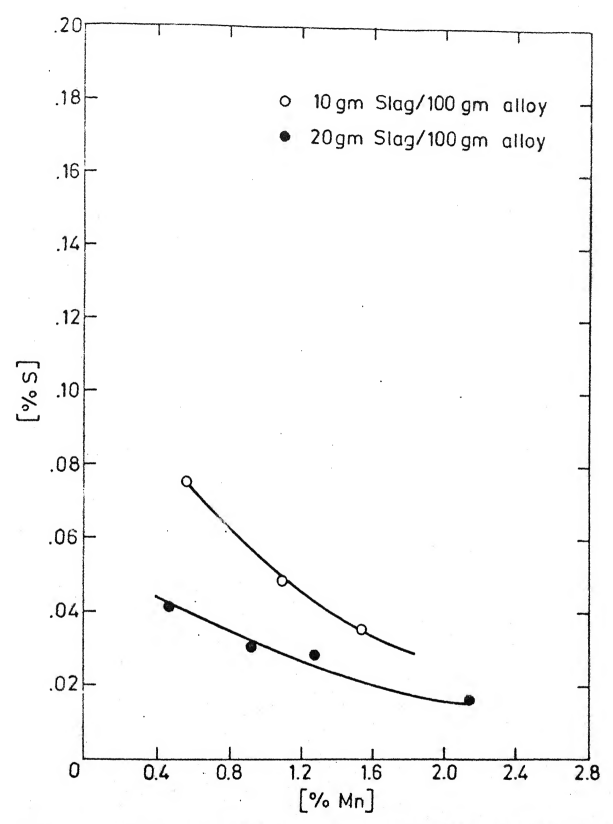


FIG. 7.7 EFFECT OF MANGANESE IN CARBON SATURATED IRON ON THE DESULPHURISATION OF THE MELT AT 1400°C AND SLAG BASICITY (CaO/SiO2 Ratio) OF 1.5.

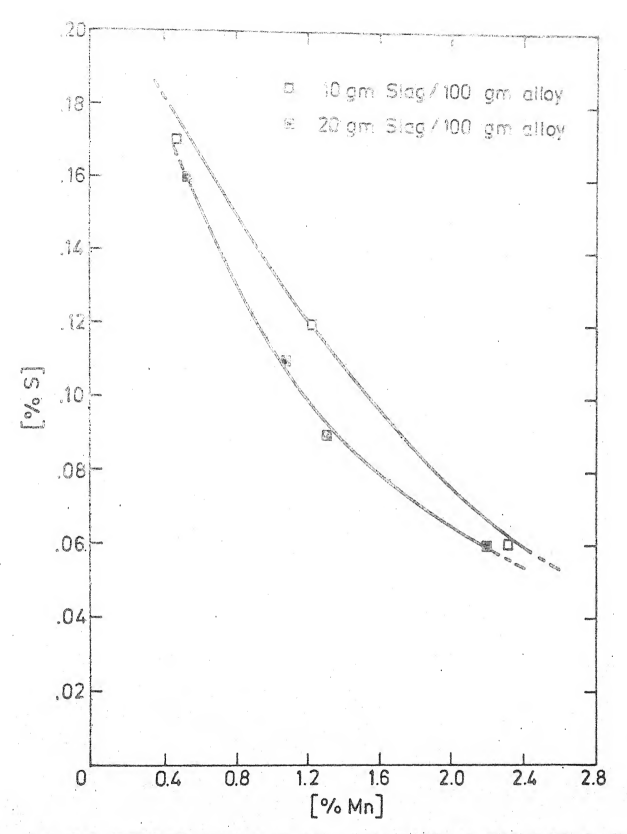


FIG. 7.8 EFFECT OF MANGANESE IN CARBON SATURATED IRON ON THE DESULPHURISATION OF THE MELT AT 1400°C AND SLAG BASICITY (CaO/SiO2 Ratio) OF 2.0.

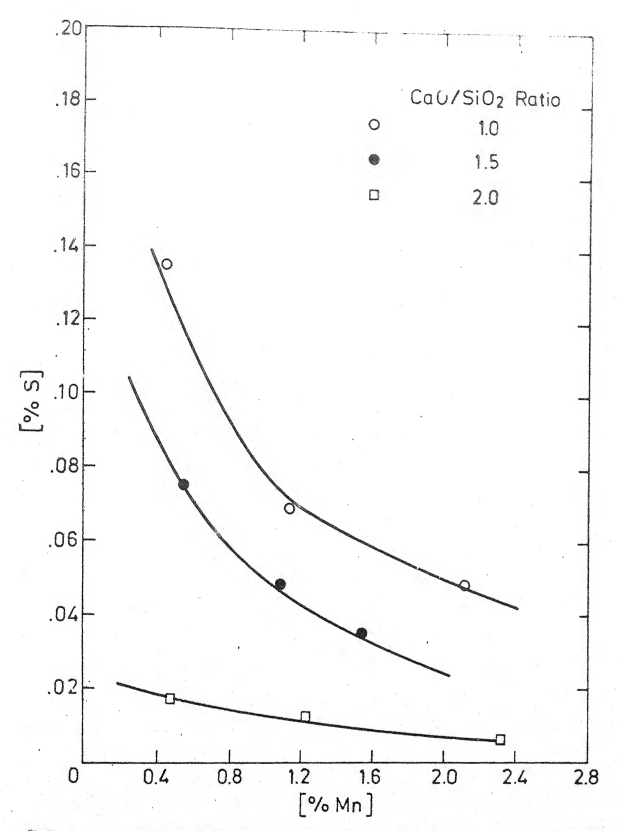


FIG. 7.9 EFFECT OF MANGANESE IN CARBON SATURATED IRON ON DESULPHURISATION OF THE MELT, CONTAINING 10 gm SLAG/100 gm OF ALLOY, AT 1400°C.

(3) Manganese in melt reacts with SiO<sub>2</sub> in slag to form silicon in the melt and MnO in slag as per reaction (2.34). This would increase slag basicity and hence favor sulphur removal.

It is possible all the three factors may be responsible for the lowering of sulphur level in presence of manganese in the melt. The probable reactions occurring in the system are further discussed in the following section.

# 7.2.3 Multi component-multiphase model

The probable reactions in the multi component system can be described by reactions (7.12) to (7.17) which are rewritten in Table 7.2. The standard free energies and equilibrium constants of these reaction can be determined from the thermodynamic data given in Chapter 2 and are included in Table 7.2.

At slag metal interface  $p_{CO}$  can be assumed to be one atm. If silicon and manganese levels are at equilibrium with respect to MnO and  $SiO_2$  in the slag by reactions (7.12) and (7.13) respectively then reaction (7.15) can describe the over all sulphur transfer from metal to slag. If reduction of oxides by carbon are ignored, silicon content in the metal will be given by reaction (7.14) and sulphur transfer can be described by reactions (7.16) or (7.17). If the reduction of  $SiO_2$  by manganese is sluggish the sulphur transfer can be described by reaction (7.12) only. If  $SiO_2$  in slag is reduced by carbon to give silicon in metal then

Table 7.2

Summary of the Reactions for Multicomponent Multiphase Model

Sl. No.	Reaction	Equa- tion No.	△G <sup>o</sup> Cal.
1	(SiO <sub>2</sub> )+2C(gr)=[Si] <sub>Wt%</sub> + 2 CO(g)	(7.12)	145100 -95.55T
2	$(MnO) + C(gr) = [Mn]_{Wt\%} + CO(g)$	(7.13)	68962.8-49.80T
3	(SiO <sub>2</sub> )+ 2[Mn] <sub>Wt</sub> =[Si] <sub>Wt</sub> +2(MnO)	(7.14)	7175 + 4.06T
4	(CaO) +[S] <sub>Wt%</sub> + C(gr) = CaS(s)+CO(g	)(7.15)	27465.7-27.12T
5	(CaO)+[S] <sub>Wt%</sub> +[Mn] <sub>Wt%</sub> =(CaS)+(MnO)	(7.16)	-41495.6+22.62T
6	$(\text{CaO})+[\text{S}]_{\text{Wt}\%}+\frac{1}{2}[\text{Si}]_{\text{Wt}\%}=(\text{CaS})+\frac{1}{2}(\text{SiO}_2)$	(7.17)	-45083 +20.65T

$$K_{7.12} = \frac{f_{si} [Si] p_{co}}{\gamma_{SiO_2} X_{SiO_2} a_c}$$
 (7.18)

Taking  $f_{Si} = 15$  and  $p_{CO} = a_{C} = 1$ , and replacing the mole fraction of  $SiO_2$  by its weight fraction, then equation (7.18) will be simplified to the following form;

$$K_{7.12} = \frac{1500 [Si]}{\gamma_{Si0_2} (\%t \% Si0_2)}$$
 (7.19)

The equilibrium sulphur level in this case will be governed by the following equation;

$$\frac{\text{(S)}}{\text{[S]}} = 4 \left(\frac{\gamma_{\text{CaO}}}{\gamma_{\text{CaS}}}\right) \quad \text{(Wt\% CaO)} \quad \text{K}_{\text{S}}$$
 (7.20)

 $\gamma_{SiO_2}$  and  $\gamma_{CaS}$  are known to vary with slag basicities as mentioned in Chapter 2. As reduction of  $SiO_2$  of carbon proceeds, slag basicity will increase which will lower  $\gamma_{SiO_2}$  and increase  $\gamma_{CaO}$  ratio. Equilibrium between slag and metal will be reached at a particular basicity greater than the initial basicity of the slag. If 'f' is weight fraction of slag added to the melt having basicity of  $\beta_1$  with 20 pct. Al $\beta_2$  and x gms of  $\beta_2$  react with carbon to form  $\beta_1$  in 100 gms of melt, then;

$$si = \frac{28}{60} x , gm$$
 (7.21)

SiO<sub>2</sub> left in the slag = 
$$\frac{80 \text{ f}}{1 + 3_1} - x$$
, gm (7.22)

Ca0 in slag = 80 f 
$$\frac{B_1}{1 + B_1}$$
, gm (7.23)

Total slag after SiO2 reduction

$$= 100 f - x$$
 (7.24)

The CaO/SiO2 ratio can now be written as follows;

$$\frac{\text{CaO}}{\text{SiO}_2} = \frac{80 \text{ f} \frac{\text{B}_1}{1 + \text{B}_1}}{80 \text{ f} \text{ B}_1}$$

$$\frac{\text{30 f B}_1}{1 + \text{B}_1} - x$$

$$(7.25)$$

Equation (7.25) can be simplified to:

$$\frac{\text{CaO}}{\text{SiO}_2} = \frac{1}{1 - \frac{(1+\text{B1})x}{80 \text{ f B1}}}$$
 (7.26)  
Knowing  $\gamma_{\text{SiO}_2}$  as function of  $\frac{\text{CaO}}{\text{SiO}_2}$ , value of x can be

Knowing  $\gamma_{\rm SiO_2}$  as function of  $\overline{\rm SiO_2}$ , value of x can be determined for equation (7.18). Calculations show that slag volume of 20 pct and basicity of 1.0, 1.21 gms of  ${\rm SiO_2}$  will be reduced to give 0.56 pct. Silicon in metal and final  ${\rm CaO/SiO_2}$  ratio 1.2 in the slag. At slag basicity of 1.2,  $\frac{{\rm (S)}}{{\rm [S]}}$  ratio is calculated to be around 80 at 1400°C from equation (7.20) which is plotted in Figure 2.6. Corresponding sulphur level in melt should be 0.025 for slag volume of 20 pct as calculated from equation (1.2). The experimental values of sulphur level in metal obtained for this combination of slag volume and composition were much greater than the equilibrium value of 0.025 pct. The equilibrium sulphur is not influenced by the addition of manganese to the melt. The equilibrium constant of  ${\rm SiO_2}$  reduction by manganese in the melt (reaction 7.14) is given as follows;

$$\gamma_{\text{MnO}} \left( \frac{\text{Wt\% MnO}}{110} \right) 15 \text{ [Si]}$$

$$K_{7.14} = \frac{1}{0.8 \text{ [Mn]}^2 \gamma_{\text{SiO}_2} (\text{Wt\% SiO}_2)} \times 100 \quad (7.27)$$

[ Mn] / (Mn0) ratio is calculated to be around 1.0 at slag basicity 1.2 and 1400°C. For 1.2 pct manganese in metal Mn0 in slag will be around 1 to 2 pct and it can not thus influence the reactions (7.12) and (7.16) much due to corresponding changes in values of  $\gamma_{\rm SiO_2}$ , (% SiO<sub>2</sub>),  $\gamma_{\rm CaO}/\gamma_{\rm CaS}$  or (% CaS) in the slag. Manganese may thus be helping to enhance the rate of either silica reduction by carbon or the sulphur transfer from the metal to slag towards the equilibrium This aspect has not been fully studied in the present work but it is in agreement with the findings of Turkdogan and other investigators. At higher slag basicity of 1.5 and 2.0 and with pure lime as desulphuriser the equilibrium sulphur levels will be also much lower than the values obtained experimentally. There also manganese may be helping in only increasing the rate of sulphur transfer from metal to the slag. The formation of MnS by the reaction between manganese and sulphur is also unlikely because the experimental data correspond to  $a_{MnS} = 0.30$  at slag basicity 1.0 and slag volume of 20 pct. Total MnS formed by the addition of manganese to the melt can not be greater than 1.0 pct.of the slag. More over  $a_{MnS}$  values calculated from the data of manganese and sulphur in the melt does not increase with increase in manganese content of metal which can be expected if MnS is formed to lower sulphur in metal. Activities of MnS in slag are

calculated to be 0.15 and 0.05 at slag basicities of 1.5 and 2.0. For manganese and sulphur reaction to occur, MnS must lower its activity by its immediate solution with slag. This will require much greater stirring than obtained in the present study.

It may also be possible that experimental results are subject to error due to improper sampling in presence of large quantity of slag. To check this samples were collected at the end of experiment from the solidified mass in the crucible. The experimental results in Table 6.6 confirm that equilibrium conditions are not reached in such experiments also. Hence manganese may be helping just to increase the rate of sulphur transfer from metal to slag phase. This aspect however is not confirmed by the results of heats H<sub>28</sub> and H<sub>29</sub> in Table 6.5. Therefore the role of Mn on the desulphurisation of the iron - carbon melt by the oxide slags of CaO-SiO<sub>2</sub>-MnO is not well understood yet and further work is needed in this area.

#### 7.3 Industrial Applications

Equilibrium sulphur level of metal in presence of manganese decreases rapidly with increase in manganese content of metal or decrease in temperature. At 1300°C and 2 pct. manganese, equilibrium sulphur value is around 0.058 pct. In the laboratory near equilibrium value i.e. 0.048 is reached in 10 minutes of cooling from 1400°C without stirring the bath. Cooling to 1200°C in 5 minutes decreases sulphur content to 0.044 pct. Thus reaction between manganese

and sulphur to form MnS product is quite fast and one can expect significant drop in sulphur content of metal of the ladle in cooling slowly during the transit from blast furnace to mixer. Data from Rourkela Steel Plant clearly shows this trend of lowering of sulphur content of metal (Figure 2.11). Presence of blast furnace carry over slag with the metal can not be ruled out and it may favor sulphur removal by manganese to form MnS as activity of MnS may be lowered. The kinetics of sulphur transfer to the oxide slag in presence of manganese without stirring the melt has not been yet studied but the equilibrium walue of sulphur is much lower than found experimentally in the laboratory condition or found in the plant data.

Iyengar et al<sup>40</sup> reported that Rourkela Steel Plant developed a norm of hot metal manganese of 1.8 pct. for obtaining bath (steel) sulphur of around 0.030 pct. Sulphur in mixer metal would be around 0.035 pct. Equilibrium data shows that temperatures for this combination of manganese and sulphur in the hot metal is around 1220°C which may be very close to the actual temperature of mixer metal though no measurements have been reported. The metal depth in the ladle or in the mixer may be 2-3 m. and in absence of stirring pure MnS must be formed by the reaction between manganese and sulphur inside the melt. Only at the slag metal interface MnS can lower its activity by its solution in slag. Diffusion of manganese and sulphur to the slag metal interface will be very slow to account for any significant drop in sulphur. Further drop may be possible only with good stirring of the melt.

#### CHAPTER VIII

#### SUMMARY AND CONCLUSIONS

Studies on sulphur control in iron making and steel making processes or by means of external treatment of metal in recent years have gained importance to meet challanges for production of low sulphur steel at lower cost for newer and newer applications in space, ocean, defence etc. Thermodynamics of sulphur reactions and desulphurising powers of various blast furnace and steel making slags have been investigated by many workers by equilibrating the melt with known  $\rm H_2/H_2S$  ratio or with a known quantity and composition of slag at a fixed temperature. In the present work attempts have been made to investigate the effect of manganese on the desulphurisation of iron—carbon melts due to formation of MnS with and without addition of some oxide slags.

Theoretical models have been proposed to calculate the sulphur content of metal as function of temperature and manganese content of metal from the reported thermodynamic data on standard free energies of formation and heats of solution. Values of activity coefficients of sulphur in iron-carbon alloys in temperature range of 1200° to 1400° C have been calculated, using a regular solution model, from the reported data at 1600° C. The experimental results of many investigation such as Oelsen, Sherman and Chipman, Morris etc. agreed with results of theoretical calculations. For sulphur

levels greater than 0.10 pct, both FeS and MnS may be formed in the slag whereas at lower values of sulphur, MnS is the predominant phase of the slag. In the present work FeS and MnS are assumed to behave ideally.

The experiments were conducted in a silicon carbide furnace under the neutral atmosphere of argon. The charge materials were put in a graphite crucible for melting in the furnace.

The calibrated thermocouples were used to measure the temperature of the crucible just above the melt accurately. Drawing alloy samples by quartz tube did not pose any problem. In a few cases samples were taken from the solidified alloy mass at the end of the experiment.

The set up for sulphur analysis of the alloy was calibrated with standard steel samples obtained from BCS and hence the results of analysis are considered to be quite accurate. The purity of the charge materials was checked by the X-ray diffraction and chemical analysis. All materials except locally procured FeS and MnS were found to be free of impurities.

Equilibrium in the Fe-C-Mn-S system was checked by drawing samples at about 1300°C both during cooling and heating cycles. The equilibrium was more or less reached in one hour duration at constant temperature.

Additional experiments were carried out to study the approach to the equilibrium during cooling of the melt without any stirring as to simulate the conditions in the ladle during

transit from the blast furnace section to the steel making section. Near equilibrium values of manganese and sulphur levels in melt were obtained in such experiments. To check whether MnS is formed in the experiments, one slag sample was chemically analysed to contain high manganese and sulphur. This was further contirmed by the presence of MnS phase in the X-ray diffraction pattern.

In some experiments manganese was added to the melt in steps at a fixed temperature. The lower content of the manganese in the melt than that of the charge suggested that maganese had reacted with sulphur in the melt to form MnS. However the formation of FeS in the slag at higher temperatures could not be ruled out.

Comparison of theoritical values of sulphur contents in carbon saturated metal at different temperature and manganese content of metal with the experimental results of present work shows that good agreement is reached in most of the cases if the activities of MnS and FeS in slag are taken same as their weight fractions and a discrepancy of - 1.0 to -1.8 KCal/mole in the reported standard free energy of formation of MnS is allowed. These limits are within the accuracy of the reported data.

From slag metal equilibrium studies for 4-6 hours at 1400°C, it is found that manganese up to 2 pct. decreases the sulphur content of metal at all combinations of slag volume (10 to 20 pct.) and slag basicity (Cao/SiO<sub>2</sub> ratios, 1 to 2).

A multicomponent multiphase model is proposed to study the approach of the system towards equilibrium. Reduction of SiO2 in the slag by carbon or manganese in the melt occurs, increasing there by the basicity of slag. For a melt having slag volume of 20 pct. and basicity of 1.0, 1.21 gms of  $SiO_2$  is reduced to give 0.56 pct. of Si in metal and  $\frac{\text{CaO}}{\text{SiO}_2}$  ratio of 1.2 in the slag. For this slag basicity, the sulphur equilibrium partition ratio is around 80 and equilibrium sulphur content is 0.025 pct. The actual values are higher than this but they tend to reach equilibrium value with increasing manganese content of metal. Equilibrium and actual sulphur content of metal decreases with increase in slag volume or increase in slag basicity. The exact role of manganese in lowering of sulphur level in the melt in presence of oxide slag however could not be properly understood from the findings and experimental results of this work. More experiments and complete analysis of metal and slag samples are needed and recommended to gain further understanding of Mn-S interactions in the Fe-C alloys.

#### CHAPTER IX

#### SUGGESTIONS FOR FURTHER WORK

Difficulties were faced for procurement of pure manganese sulphide material. The manganese sulphide procured locally contained lot of unidentified impurities. Hence it is suggested that attempts should be made for preparation of manganese sulphide and iron sulphide in the laboratory by some suitable methods such as synthesising pure manganese and pure sulphur under vacuum around 250°C.

Basic studies by addition of MnS to iron-carbon melt may be necessary to understand its decomposition behavior in the melt and give an independent confirmation of manganese sulphur equilibrium in the Fe-C-Mn-S melt.

In order to explain the effect of manganese on desulphurisation in presence of oxide slags, synthetic slags such as MnS-CaO-SiO<sub>2</sub>-Al<sub>2</sub>O<sub>3</sub> and FeS-CaO-SiO<sub>2</sub>-Al<sub>2</sub>O<sub>3</sub> should be prepared in the laboratory for studying slagmetal equilibrium.

Further experiments may be planned for equilib-rating the Fe-C-Mn melts with known  $\rm H_2/\rm H_2S$  ratio at fixed temperatures.  $\rm H_2/\rm H_2S$  ratio of  $\rm H_2$  and  $\rm H_2S$  gas mixture will correspond to a particular sulphur concentration in the melt at equilibrium and this in turn will correspond to a particular manganese content of metal. The final manganese content of the alloy melt can thus be compared with the theoretical value of

manganese-sulphur equilibrium calculations. The starting manganese in the melt must be kept greater than the equilibrium value for such experiments.

It is desirable to have access to complete and accurate analysis of slag and metal for better correlations of the thermodynamic data and comparison with predicted values.

Experimental data is also needed for melts containing carbon other than saturation values. This may be possible in the pure  ${\rm Al}_2{}^0{}_3$  cyacibles.

There is also need to get more scientific and reliable data from the industry.  $\cdot$ 

Temperature of blast furnace metal and slag, accurate analysis of blast furnace metal and slag, ladle metal temperature and analysis, mixer metal and mixer slag analysis, mixer temperature are needed to correlate the industrial data with laboratory study. Industrial data of desulphurisation of iron with additions of pure lime, slags and carbides in presence of manganese in the metal will also be helpful in analysis of data.

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